	I.1. Consignor				I.2. IMSOC Reference				
	Name				I.2.a. Local Reference				
	Address								
	Country		ISO Code						
	I.5. Consignee				I.3. Central competent authority				
H	Name				I.4. Local comp	etent authority			
ler	Address				1.1. Locar comp	cicili dudiority			
Ы	Country		ISO Code						
Part I : Details of consignment	I.7. Country of ori	gin		ISO Code	I.9. Country of destination ISO Code			ISO Code	
É C	I.8. Region of orig	in		Code	I.10. Region of (destination			
S 0	I.11. Place of Disp				I.12. Place of de				
tai	Name				Name				
Dei	Address				Address				
•••	Approval Numbe	r			Approval Num	ber			
ť	Country		ISO Code		Country		ISO Code		
Pai									
	I.13. Place of Load	ling			I.14. Date and t	ime of departure			
	Name								
	Address								
	Approval Numbe	er							
	Country		ISO Code						
	I.15. Means of Tra	insport			I.16 Entry Point				
	Mode	International	Identification						
		transport document							
					-				
					-				
					_				
	I.18. Transport co		_	_	I.17. Accompanying documents				
	Frozen 🗖	Controlled temperature 🛛	Ambient 🛛	Chilled \Box	Accompanying document reference Date of issue Country				
					Place of issue				
	I.19. Container No	o / Seal No							
	I.20. Certified as								
	Pet food 🛛		Technical use 🛛						
		rough a third cour							
	1.21. For transit in	irough a third cour	itry		I.22. For transit through Member State(s)				
	Country		ISO Code		Country	IS	SO Code		
	EU Exit Authority		BCP code		-				
	EU Entry		BCP code						
	Authority				-				
	I.25. Total net wei	ght			I.25. Total gross weight				
	I.28. Description o	of consignment							
	1. 35 ALBUMINOI	DAL SUBSTANCES;	MODIFIED STARC	HES; GLUES; ENZY	MES				
		seinates and other							
	#1. Commodity		pecies	Manufactur	ring plant	Net weight	Batch nur	nber	
			•						

	II. Health info	rmation								
	the Europe Regulation	ean Parliam (EU) No 14	ent and of th	ne Council, a in particular	re that I have read and understood Regulation (EC) No 1069/2009 of and in particular Articles 8 and 10 thereof, and Commission r Chapter II of Annex XIII and Chapter II of Annex XIV thereto, and					
ion	II.1.				plant approved and supervised by the competent authority in alation (EC) No 1069/2009;					
icat	II.2.	has been p	orepared exc	clusively with	h the following animal by-products:					
Part II: Certification		(2)	either 🗆 [-	of animals	nd parts of animals slaughtered or, in the case of game, bodies or parts killed, and which are fit for human consumption in accordance with U law, but are not intended for human consumption for commercial					
Pa		(2)	and/or □ [-	slaughtered human con following p	carcases and the following parts originating either from animals that have been slaughtered in a slaughterhouse and were considered fit for slaughter for human consumption following an ante-mortem inspection or bodies and the following parts of animals from game killed for human consumption in accordance with retained EU law:					
	-			(i)	carcases or bodies and parts of animals which are rejected as unfit for human consumption in accordance with retained EU law, but which did not show any signs of disease communicable to humans or animals;					
				(ii)	heads of poultry;					
				(iii)	hides and skins, including trimmings and splitting thereof, horns and feet, including the phalanges and the carpus and metacarpus bones, tarsus and metatarsus bones;					
				(iv)	pig bristles;					
				(v)	feathers;]					
		(2)	and/or □ [-	referred to Parliament	products from poultry and lagomorphs slaughtered on the farm as o in Article 1(3)(d) of Regulation (EC) No 853/2004 of the European t and of the Council, which did not show any signs of disease able to humans or animals]					
		(2)	and/or □ [-	through blo slaughtered for human	nimals which did not show any signs of disease communicable ood to humans or animals, obtained from animals that have been d in a slaughterhouse after having been considered fit for slaughter consumption following an ante-mortem inspection in accordance ned EU law;]					
		(2)	and/or □ [-	consumptio	products arising from the production of products intended for human on, including degreased bone, greaves and centrifuge or separator m milk processing;]					
		(2)	and/or □ [-	which are r or due to p	f animal origin, or foodstuffs containing products of animal origin, no longer intended for human consumption for commercial reasons problems of manufacturing or packaging defects or other defects from risk to public or animal health arise;]					
		(2)	and/or □ [-	by-product commercia	Id feedingstuffs of animal origin, or feedingstuffs containing animal ts or derived products, which are no longer intended for feeding for al reasons or due to problems of manufacturing or packaging defects efects from which no risk to public or animal health arise;]					
		(2)	and/or □ [-	from live a	centa, wool, feathers, hair, horns, hoof cuts and raw milk originating nnimals that did not show signs of any disease communicable through ct to humans or animals;]					
		(2)	and/or □ [-		imals, and parts of such animals, except sea mammals, which did not signs of diseases communicable to humans or animals;]					
		(2)	and/or □ [-		products from aquatic animals originating from plants or tents manufacturing products for human consumption;]					

	II. Health information							
	(2)	and/or □ [-	the following material originating from animals which did not show any signs of disease communicable through that material to humans or animals:					
			(i) shells from shellfish with soft tissue or flesh;					
			(ii)	the follow	ing originat	ing from terrestria	l animals:	
l g				_	hatchery b	oy-products,		
atic				_	eggs,			
lfic				_	egg by-pro	ducts, including eg	gg shells,	
l T			(iii)	day-old ch	icks killed f	or commercial rea	sons;]	
[- except Category 1 material as r							tebrates other than species	
					parts thereof of the zoological orders of Rodentia and Lagomorpha, ry 1 material as referred to in Article 8(a)(iii), (iv) and (v) of C) No 1069/2009 and Category 2 material as referred to in Article			
	(2)	and/or □ [-	are prohik	oited by Cou	ncil Directiv	ve 96/22/EC, the imj	h certain substances which port of the material being ulation (EC) No 1069/2009;]	
	II.3.		1	1				
	(2)						ughout its substance;]	
	(2)	or o lwas	produced as regards ingredients of animal origin using exclusively products been:					
			(a)	in the case of animal by-products or derived products from mea meat products subjected to a heat treatment of at least 90 °C throughout its substance;				
				e of milk and	d milk based produ	icts,		
				(i)	countries Regulation	listed in column B n (EU) No 605/2010 ition treatment suf	ies or parts of third of Annex I to Commission submitted to a ficient to produce a negative	
				(ii)	parts of th Regulation	ird countries listed n (EU) No 605/2010, ition treatment suf	n 6 from third countries or l in column C of Annex I to , first submitted to a ficient to produce a negative	
				(iii)	countries No 605/20 double he	listed in column C 10, submitted to a s at treatment where	ies or parts of third of Annex I to Regulation (EU) sterilisation process or a e each treatment was ive phosphatase test on its	
				(iv)	countries No 605/20 and-mout vaccinatio carried ou	listed in column C (10, where there ha h disease in the pre n against foot-and	ies or parts of third of Annex I to Regulation (EU) s been an outbreak of foot- eceding 12 months or where -mouth disease has been 12 months, submitted to	
					either —		ocess whereby an Fc value	
					or	equal of greater	than 3 is achieved	

	II. Health information					
u			at pa le ne	least equal to t asteurisation pr ast 15 seconds	eatment with a heating that achieved by a cocess of at least 72 °C and sufficient to produ n to a phosphatase test	for at uce a
cati(either			
Part II: Certification			at he su ph dr	least equal to t eat treatment, a ifficient to proc nosphatase test	eatment with a heating that achieved by the ir and which would be duce a negative reaction , followed, in the case ied milk-based produce	nitial on to a of
			or			
			be		process such that the p l at less than 6 for at le	
	(c)	in the case of gelatine, produced using a process that ensure unprocessed Category 3 material is subjected to a treatmen acid or alkali, followed by one or more rinses with subsequ adjustment of the pH and subsequent, if necessary repeated extraction by heat, followed by purification by means of filt and sterilisation;				rith t
	(d)	in the case of hydrolysed protein produced using a production process involving appropriate measures to minimise contamination of raw Category 3 material, and, in the case of hydrolysed protein entirely or partly derived from ruminant hides and skins produced in a processing plant dedicated only to hydrolysed protein production, using only material with a molecular weight below 10000 Dalton and a process involving the preparation of raw Category 3 material by brining, liming and intensive washing followed by:				nation tein luced
		 (i) exposure of the material to a pH of more than 11 more than three hours at a temperature of more to °C and subsequently by heat treatment at more the °C for 30 minutes at more than 3,6 bar; or 		emperature of more th treatment at more tha	nan 80	
		(ii)		an 11, followed	n pH of 1 to 2, followed d by heat treatment at	
	(e)	methods 1 Regulatior	to 5 or 7, as ref (EU) No 142/20	ferred to in Cha 011; or treated :	any of the processing apter III of Annex IV to in accordance with Ch (EC) No 853/2004 ;	
	(f)	in the case of collagen submitted to a process ensuring the unprocessed Category 3 material is subjected to a treatme involving washing, pH adjustment using acid or alkali foll one or more rinses, filtration and extrusion, the use of pro- other than those permitted by retained EU law being pro-				vatives
	(g)	g) in the case of blood products, produced using any methods 1 to 5 or 7, as referred to in Chapter III of Regulation (EU) No 142/2011;			-	
	(h)	in the case of mammalian processed animal protein submitted any of the processing methods 1 to 5 or 7 and, in the case of po blood, submitted to any of the processing methods 1 to 5 or 7				

Part II: Certification

ROPEAN UNION II. Health information	((50)11000	essed petfood from EU countries 3B 142-2011 GBHC092E			
		-	d that in the case of method 7 a heat treatment throughout its ce at a minimum temperature of 80 °C has been applied;			
	(i)	of fishn	ase of non-mammalian processed protein with the exclusion neal submitted to any of the processing methods 1 to 5 or 7 as d to in Chapter III of Annex IV to Regulation (EU) No 142/2011;			
	(j)	to 7 as r 142/201 product	ase of fishmeal submitted to any of the processing methods 1 referred to in Chapter III of Annex IV to Regulation (EU) No 1 or to a method and parameters which ensure that the complies with the microbiological standards for derived as set out in Chapter I of Annex X to Regulation (EU) No 1;			
	(k)	the proc oil) as r 142/201 Annex l rumina	ase of rendered fat, including fish oils, submitted to any of cessing methods 1 to 5 or 7 (and method 6 in the case of fish eferred to in Chapter III of Annex IV to Regulation (EU) No 1 or produced in accordance with Chapter II of Section XII of II to Regulation (EC) No 853/2004; rendered fats from nt animals must be purified in such a way that the maximum the remaining total insoluble impurities does not excess 0,15 ight;			
	(1)	in the c	ase of dicalcium phosphate produced by a process that			
		(i)	ensures that all Category 3 bone-material is finely crushed and degreased with hot water and treated with dilute hydrochloric acid (at a minimum concentration of 4 % and a pH of less than 1,5) over a period of at least two days;			
		(ii)	following the procedure referred to in (i), applies a treatment of the obtained phosphoric liquor with lime, resulting in a precipitate of dicalcium phosphate at pH 4 to 7; and			
		(iii)	finally, air dries the precipitate of dicalcium phosphate with inlet temperature of 65 °C to 325 °C and end temperature between 30 °C and 65 °C ;			
	(m)	in the ca ensures	ase of tricalcium phosphate produced by a process that			
		(i)	that all Category 3 bone-material is finely crushed and degreased in counter-flow with hot water (bone chips less than 14 mm);			
		(ii)	continuous cooking with steam at 145 °C during 30 minutes at 4 bar;			
		(iii)	separation of the protein broth from the hydroxyapatite (tricalcium phosphate) by centrifugation; and			
		(iv)	granulation of the tricalcium phosphate after drying in a fluid bed with air at 200 °C ;			
	(n)	method	ase of flavouring innards, produced according to a treatment and parameters, which ensure that the product complies e microbiological standards referred to in point II.4.]			
(2)	or \circ [was subject to a treatment such as drying or fermentation, which has been authorised by the competent authority;]					
(2)	humans or animals, h	as been s and which	terrestrial invertebrates other than species pathogenic to ubject to a treatment which has been authorised by the a ensures that the petfood poses no unacceptable risks to			
II.4. was ana	-		east five samples from each processed batch taken during or			

ΕU	ROPEAN U	JNION		(1	JB) Processed	i petiooa irom i	EU countr	1es 3B 142-2011 GBHC09			
	II. Health info	rmation									
		after stora	orage at the processing plant and complies with the following standards (4):								
		after storage at the processing plant and complies with the following standards (4): Salmonella: absence in 25g: n = 5, c = 0, m = 0, M = 0,									
					0	10, M = 300 in 1	ramma				
	II.5.	hasundar						nte ofter treatment:			
d						-	• •	nts after treatment;			
Part II: Certification	II.6.	it is clearly	red in new packaging, Which, if the petfood is not dispatched in ready-to-sell packages on which ly indicated that the content is destined for feeding to pets only, bear labels indicating "NOT MAN CONSUMPTION";								
erti	(2) 🗆 [II.7.	the petfoo	described above								
С Н		(2)	either \circ [is derived from other ruminants than bovine, ovine or caprine animals.]								
Part]		(2)	or ○ [is d derived f		bovine, ovine	or caprine anima	ls and does	s not contain and is not			
			(2)	animals b classified	either \circ [bovine, ovine and caprine materials other than those derived from animals born, continuously reared and slaughtered in a country or region classified as posing a negligible BSE risk in accordance with Decision 2007/453/EC.]]						
(2) or ○ [(a) specified risk material as (EC) No 999/2001 of the European							-	•			
				(b)	or caprine an continuously classified as	nimals, except fro v reared and slau posing a negligib Decision 2007/45	om those a ghtered in le BSE risk	from bones of bovine, ovin nimals that were born, a country or region in accordance with hich there has been no			
				(c)	or caprine at laceration of rod-shaped i means of gas animals that country or ro	nimals which have the central nerv nstrument introc injected into the were born, conti	ve been kill ous tissue l luced into e cranial ca inuously re s posing a l	btained from bovine, ovin led, after stunning, by by means of an elongated the cranial cavity, or by wity, except for those eared and slaughtered in a negligible BSE risk in []			
	Notes										
		ountries sul celand and S			import arrang	gements include:	an EU men	nber State; Liechtenstein;			
	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).										
References to Great Britain in this certificate include Channel Islands and Isle of Man.											
	Part I:										
 Box Person responsible for the consignment in Great Britain, Channel Islands or reference los is required to be filled in only if it is a certificate for a commodity to be through Great Britain, Channel Islands or Isle of Man; it may be filled in if for a commodity to be imported into Great Britain, Channel Islands or Isle 						nodity to be transited e filled in if the certificate i					
		Box reference I.12:	commodi					icate for a transit , free warehouses and			
		Boy	Decistrat	ion numbor	(noilusou uso go	aa an aantainan a	nd lormioa)	flight number (aircraft) o			

	II. Health info	rmation								
		I.16:								
	_	Box reference I.19:		ystem (HS) code under the following headings: 04.01; .05, 05.06; 05.11, 15.01, 15.02, 15.03, 15.04, 23.01, 23.09; 03 or 35.04.						
cation	_	Box reference I.23:	for bulk containers; the container given.	number and the seal number	(if applicable) must be					
Part II: Certification	_	Box reference I.25:	technical use: any use other than f production or manufacturing of pe	feeding of farmed animals, other than fur animals, and the et food.						
Part I	_	Box reference I.26 and I.27:	fill in according to whether it is a transit or an import certificate.							
	_	Box reference I.28:	Species: select from the following: Aves, Ruminantia, Suidae, Mammalia other than Ruminantia or Suidae, Pesca, Mollusca, Crustacea, Invertebrates other than Mollusca crustacea.							
	PART II:									
	(2)	Delete as a	ppropriate.							
	(4)	Where:								
			n = number of samples to be tested	d;						
			m = threshold value for the number number of bacteria in all samples		isidered satisfactory if the					
			M = maximum value for the numb the number of bacteria in one or n		-					
			-	al count of which may be between m and M, the sample The bacterial count of the other samples is m or less.						
	_	The signatu	are and the stamp must be in a diffe	erent colour to that of the printing.						
	_	This certifi	ne person responsible for the consignment in Great Britain, Channel Islands or Isle of Man: īcate is only for veterinary purposes and must accompany the consignment until it reaches r control post of entry into Great Britain, Channel Islands or Isle of Man.							
	Certifying Offi									
	Name (in capit Date of signati		Qualification and title Signature							
	Stamp	ure		Signature						