Official certificate to the EU

	I.1. Consignor					I.2. Certificate	e reference	I.2.a.	IMSOC r	eference		
	Name					I.3. Central competent authority Specimen not to be used for imports into			to the			
	Address					I.4. Local competent authority						
	Country			ISO Code								
İ	I.5. Consignee					I.6. Responsib	le for the consigni	nent in	EU			
1	Name					Name	o .					
en	Address					Address						
틹	Country			ISO Code		Country			ISO C	ode		
: Details of consignment	I.7. Country of ori	gin	ISO Code	I.8. Region of origi	in Code	I.9. Country of destination	of ISO C	code I	.10. Regio	on of destinati	on	Code
8	I.11. Place of dispa	ıtch				I.12. Place of	doctination					
oĮ	_	itti				Name	uestination					
ils	Name Address					Address						
eta	Approval Number	r				Approval Nu	mber					
Ă	Country	-		ISO Code		Country	11.201		IS	O Code		
H١												
Part	I.13. Place of loadi	ng				I.14. Date and	l time of departure	!				
ŭ	Name											
	Address											
	Approval Number	r										
	Country			ISO Code								
ł	I.15. Means of Trai	nsnort				I.16. Entry po	int					
	Mode	-	ational	Identification		1.10. Entry po						
	Mode	transp	ort	identification								
		docun	nent									
ŀ	I.18. Transport cor	nditions	,			I 17 Accomps	anying documents					
	Frozen		Chilled 🗆	Ambie	ent 🗆	Type Number	mynig documents					
İ	I.19. Container No	/ Seal N	Io		7							
		, ocar i	NU)) '							
•	I.20. Certified as				, \ \ '							
•	I.20. Certified as			d Slau Petfo Oth	e Fatte Furt Ro	egis Cann Gar	n Feed Appr M	an Br	ee Pets	Prod Trad	Rela	Furt
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	I.20. Certified as Trav Phar Quar ellin mace antin g utica e circu use estab s/ani mal ment acts I.21. For transit	Tech nical Use	Artifi Procial uct repr for oduc hution an ucor	n is pt	ing n	I.22. For inter	es of pe	tfo n	ŤÍO	uctio e n of sam petfo ples od	ying area/ purif icati on centr e	her proc essin g
	I.20. Certified as Trav Phar Quar ellin mace antin g utica e circu use estab s/ani mal acts I.21. For transit Non-EU I.24. Total number	Tech nical Use	Artifi Procial uct repr for hw tion an cor um ion	n is pt ISO Code	ing n	I.22. For inter	es of pe	tfo n	ŤÍO	uctio e n of sam petfo ples od	ying area/ purif icati on centr e	her proc essin g
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	II. Health info	rmation									
	I, the undersigned official veterinarian, declare that I have read and understood Regulation (EC) No 1069/2009 of the European Parliament and of the Council (1a), and in particular Articles 8 and 10 thereof, and Commission Regulation (EU) No 142/2011 (1b), and in particular Chapter II of Annex XIII and Chapter II of Annex XIV thereto, and certify that the petfood described above:										
	II.1. has been prepared and stored in a plant approved and supervised by the competent authority in accordance with Article 24 of Regulation (EC) No 1069/2009;										
ati	II.2.	2. has been prepared exclusively with the following animal by-products:									
Part II : Certification		(2)	□ either	[-	bodies or p	nd parts of animals slaughtere arts of animals killed, and wh on in accordance with Union l or human consumption for co	nich are fit for human legislation, but are not				
Par		(2)	□ and/or	[-	have been s for slaughte inspection	nd the following parts original slaughtered in a slaughterhou er for human consumption fo or bodies and the following p uman consumption in accord	use and were considered fit llowing an ante-mortem arts of animals from game				
					(i)	carcases or bodies and parts rejected as unfit for human of with Union legislation, but w of disease communicable to	consumption in accordance which did not show any signs				
					(ii)	heads of poultry;					
					(iii)	hides and skins, including tri thereof, horns and feet, inclu carpus and metacarpus bone bones;	iding the phalanges and the				
					(iv)	pig bristles;					
					(v)	feathers;]					
		(2)	□ and/or	[-	farm as ref	products from poultry and lag erred to in Article 1(3)(d) of R pean Parliament and of the C igns of disease communicable	egulation (EC) No 853/2004 ouncil (2a), which did not				
		(2)	□ and/or	[-	communica animals that been consid	imals which did not show any able through blood to humans at have been slaughtered in a dered fit for slaughter for hur artem inspection in accordance	s or animals, obtained from slaughterhouse after having nan consumption following				
		(2)	□ and/or	[-	intended fo	products arising from the pro or human consumption, includ d centrifuge or separator slud	ding degreased bone,				
		(2)	□ and/or	[-	animal original for commen	rcial reasons or due to proble defects or other defects from	ded for human consumption ms of manufacturing or				
		(2)	□ and/or	[-	containing longer inter problems o	d feedingstuffs of animal orig animal by-products or derive nded for feeding for commero f manufacturing or packaging n no risk to public or animal h	d products, which are no cial reasons or due to g defects or other defects				
		(2)	□ and/or	[-	originating	enta, wool, feathers, hair, hor from live animals that did no able through that product to h	ot show signs of any disease				

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		II. Health information								
		(2)	□ and/or	[-	_	not show any signs of d		als, except sea mammals, s communicable to humans		
		(2)	□ and/or	[-	animal by-products from aquatic animals originating from pl establishments manufacturing products for human consump					
10:100	rari II : cerimicanon	(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:					
	err				(i)	shells from shellfish v	vith sof	t tissue or flesh;		
19	ا:				(ii)	the following originat	ing fro	m terrestrial animals:		
1	I.I					- hatchery l	_			
6	Ľя					- eggs,				
							ducts,	including egg shells,		
					(iii)	day-old chicks killed f		9 99		
		(2)	□ and/or	[-		•		estrial invertebrates other		
		. ,	, .	•		s pathogenic to humai				
		(2)	□ and/or	[-	Lagomorph 8(a)(iii), (iv)	a, except Category 1 n	naterial (EC) No	cal orders of Rodentia and as referred to in Article o 1069/2009 and Category 2 g) of that Regulation;]		
		(2)	□ and/or	[-	substances the import		by Cour permitte	ncil Directive 96/22/EC (2b), ed in accordance with		
		II.3.								
		(2)	o either	[was subject	ted to a hea	t treatment of at least	90°C tł	nroughout its substance;]		
		(2)	o or		ced as regar hich had be	ds ingredients of anin en:	nal orig	in using exclusively		
			C	(a)	meat produ	of animal by-products acts subjected to a heat its substance;		ved products from meat or nent of at least 90 °C		
				(b)	in the case	of milk and milk based	d produ	cts,		
						Regulation (EU) No 60	umn B 05/2010	of Annex I to Commission		
					(ii)	with a pH reduced to parts of third countrie Regulation (EU) No 60	es listed 5/2010,	in 6 from third countries or l in column C of Annex I to first submitted to a ficient to produce a negative		
					(iii)	if they are from third countries listed in col No 605/2010, submitte double heat treatmen	umn C o ed to a s t where	of Annex I to Regulation (EU) sterilisation process or a		

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	II. Health information				
uo	(iv)	countries No 605/20 and-mout vaccination carried on	if they are from third countries or parts of third countries listed in column C of Annex I to Regulation (EU No 605/2010, where there has been an outbreak of footand-mouth disease in the preceding 12 months or where vaccination against foot-and-mouth disease has been carried out in the preceding12 months, submitted to		
tificati		o either -		ocess whereby an Fc value	
Cer		o or	equal or greater	than 3 is achieved	
Part II : Certification		-	at least equal to to pasteurisation proleast 15 seconds	eatment with a heating effect that achieved by a rocess of at least 72 °C for at and sufficient to produce a n to a phosphatase test,	
			o either		
			heati that a treat suffic react follow milk,	ond heat treatment with a ng effect at least equal to achieved by the initial heat ment, and which would be cient to produce a negative ion to a phosphatase test, wed, in the case of dried or dried milk-based ucts by a drying process	
			o or	acto 27 a a1711.6 p200000	
			the p	sidification process such that H has been maintained at han 6 for at least one hour;	
	unproces acid or al adjustme	sed Category kali, followe nt of the pH n by heat, fo	y 3 material is subjeed by one or more r and subsequent, if	process that ensures that ected to a treatment with inses with subsequent necessary repeated, ion by means of filtration	

(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:

- (i) exposure of the material to a pH of more than 11 for more than three hours at a temperature of more than 80 °C and subsequently by heat treatment at more than 140 °C for 30 minutes at more than 3,6 bar; or
- (ii) exposure of the material to a pH of 1 to 2, followed by a pH of more than 11, followed by heat treatment at 140 °C for 30 minutes at 3 bar;

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	ountry	172/20	11 (2013/3)	19) 3(b) Frocesseu peu	ood other than canned petto	ou
	II. Health information					
		(e)	methods 1 Regulation	to 5 or 7, as referred to it (EU) No 142/2011; or treat	ed to any of the processing n Chapter III of Annex IV to ated in accordance with to Regulation (EC) No 853/2004	;
rtification		(f)	unprocesse involving v one or mor	washing, pH adjustment t ce rinses, filtration and e	a process ensuring that subjected to a treatment using acid or alkali followed by xtrusion, the use of preservative n legislation being prohibited;	
Part II : Certification		(g)	methods 1		uced using any of the processing n Chapter III of Annex IV to	y
Pa		(h)	of the proc blood, sub- provided th	essing methods 1 to 5 or mitted to any of the proce hat in the case of method	d animal protein submitted to a 7 and, in the case of porcine essing methods 1 to 5 or 7 I 7 a heat treatment throughout are of 80 °C has been applied;	
		(i)	of fishmea	l submitted to any of the	cessed protein with the exclusion processing methods 1 to 5 or 7 a IV to Regulation (EU) No 142/201	as
		(j)	to 7 as refe 142/2011 o product co	rred to in Chapter III of A r to a method and param mplies with the microbio	o any of the processing methods Annex IV to Regulation (EU) No leters which ensure that the ological standards for derived nex X to Regulation (EU) No	1
	Ć	(k)	processing as referred 142/2011 o Annex III t ruminant a	methods 1 to 5 or 7 (and I to in Chapter III of Anno r produced in accordanc o Regulation (EC) No 853 animals must be purified e remaining total insolub	ng fish oils, submitted to any of to method 6 in the case of fish oil; ex IV to Regulation (EU) No e with Chapter II of Section XII of /2004; rendered fats from in such a way that the maximu le impurities does not excess	of
		(I)			produced by a process that	
		C	(i)	ensures that all Categor crushed and degreased dilute hydrochloric acid	ry 3 bone-material is finely with hot water and treated with d (at a minimum concentration on an 1,5) over a period of at least	
			(ii)	treatment of the obtain	e referred to in (i), applies a ed phosphoric liquor with lime, e of dicalcium phosphate at pH	
			(iii)		cipitate of dicalcium phosphate of 65 °C to 325 °C and end 0 °C and 65 °C ;	;
		(m)	in the case ensures	of tricalcium phosphate	produced by a process that	
			(i)		e-material is finely crushed and ow with hot water (bone chips	
			(ii)	continuous cooking with minutes at 4 bar;	h steam at 145 °C during 30	

142/2011 (2019/319) 3(B) Processed petfood other than canned petfood

	untry			142/20	11 (2019/31	9) 3(B) Processed petfood	onier man canneu penoou		
	II. Health info	rmation							
					(iii)	separation of the protein bro (tricalcium phosphate) by ce			
					(iv)	granulation of the tricalcium fluid bed with air at 200 °C;	n phosphate after drying in a		
cation	(n)			(n)	method an	of flavouring innards, produ d parameters, which ensure t icrobiological standards refer	hat the product complies		
ertifi	method and parameters, which ensure that the with the microbiological standards referred to i [was subject to a treatment such as drying or fermentation, authorised by the competent authority;]						ntation, which has been		
(2) or [in the case of aquatic and terrestrial i pathogenic to humans or animals, has been authorised by the competent aut poses no unacceptable risks to public a						or animals, has been subject competent authority and wh	to a treatment which has nich ensures that the petfood		
	II.4.			andom sampling of at least five samples from each processed batch taken during processing plant and complies with the following standards (4):					
		Salmonella	a: absence ir	n 25g: n = 5,	c = 0, m = 0,	$\mathbf{M}=0,$			
		Enterobact	teriaceae: n	= 5, c = 2, m	= 10, M = 30	00 in 1 gramme;			
	II.5.	has underg	gone all pred	cautions to a	ivoid contar	nination with pathogenic age	nts after treatment;		
	II.6.	it is clearly		hat the cont		etfood is not dispatched in rea ed for feeding to pets only, be	dy-to-sell packages on which ear labels indicating "NOT		
	(2) \square [II.7.	the petfood	d described	above					
		(2)	\square either	[is derived	from other	ruminants than bovine, ovine	e or caprine animals.]		
		(2)	\square and/or	[is derived not derived		e, ovine or caprine animals a	nd does not contain and is		
			(2)	o either	animals boregion clas	rine and caprine materials oth orn, continuously reared and s sified as posing a negligible B 007/453/EC.]	slaughtered in a country or		
			(2)	o or	[(a)	specified risk material as de Regulation (EC) No 999/2001 and of the Council (5);	fined in point 1 of Annex V to of the European Parliament		
(b) mechanically separated meat obtation bovine, ovine or caprine animals, animals that were born, continuous slaughtered in a country or region negligible BSE risk in accordance to Decision 2007/453/EC (6), in which				mals, except from those tinuously reared and region classified as posing a ance with Commission					
				indigenous BSE case, (c) animal by-product or derived product obtained from bovine, ovine or caprine animals which have been killed, after stunning, by laceration of the 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, except for those animals that were born, continuously reared and slaughtered in a country or region classified as posing a negligible BSE risk in accordance with Decision 2007/453/EC.]]]					
	Notes								

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Country 142/2011 (2019/319) 3(B) Processed petfood other than canne									
	II. Health info	ormation							
	Part I:								
	Turch	Box reference I.6:	Person responsible for the consignment in the European Union: this box is required to be filled in only if it is a certificate for a commodity to be transited through the European Union; it may be filled in if the certificate is for a commodity to be imported into the European Union.						
Part II : Certification		Box Place of destination: this box is to be filled in only if it is a certificate for a tra reference commodity. Products in transit may only be stored in free zones, free warehouses.							
t II : Cert		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 border inspection post of entry into the European Union.						
Par		Box reference I.19:	for bulk containers, the container number and the seal number (if applicable) must be given.						
		Box reference I.20:	technical use: any use other than feeding of farmed animals, other than fur animals, and the production or manufacturing of pet food.						
		Box reference I.21 and I.22:	fill in according to whether it is a transit or an import certificate.						
		Box reference I.25:	use the appropriate Harmonized System (HS) code under the following headings: 04.01; 04.02; 04.03; 04.04; 04.08, 05.04, 05.05, 05.06; 05.11, 15.01, 15.02, 15.03, 15.04, 23.01, 23.09; 28.35.25; 28.35.26; 35.01; 35.02; 35.03 or 35.04.						
		Box reference I.25:	Species: select from the following: Aves, Ruminantia, Suidae, Mammalia other than Ruminantia or Suidae, Pesca, Mollusca, Crustacea, Invertebrates other than Mollusca and crustacea.						
	Part II:								
	(1a)	-	4.11.2009, p. 1.						
	(1b)	OJ L 54, 26	2.2011, p. 1.						
	(2)		ppropriate.						
	(2a)	•	0.4.2004, p. 55.						
	(2b)	•	3.5.1996, p. 3.						
	(3)	-	0.7.2010, p. 1.						
	(4)	Where:							
			r of samples to be tested;						
		bacteria in	old value for the number of bacteria; the result is considered satisfactory if the number of all samples does not exceed m;						
			aximum value for the number of bacteria; the result is considered unsatisfactory if the number of ria in one or more samples is M or more; and						
			r of samples the bacterial count of which may be between m and M, the sample still being l acceptable if the bacterial count of the other samples is m or less.						
	(5)	OJ L 147, 3	1.5.2001, p. 1.						
	(6)	OJ L 172, 3	0.6.2007, p. 84.						
	-	The signat	ure and the stamp must be in a different colour to that of the printing.						
	-	veterinary	e person responsible for the consignment in the European Union: This certificate is only for purposes and must accompany the consignment until it reaches the border inspection post of the European Union.						

			and thair carried policou
	II. Health information		
	Official veterinarian or Official inspector		
	Name (in capital letters)	Qualification and title	
	Date of signature	Signature	
	Stamp	Signature	
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Part II : Certification			
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