CHAPTER 3(B)

Health certificate

For processed petfood other than canned petfood, intended for dispatch to $\frac{1}{2}$ or for transit through(²) the European Union

COUNTRY: UNITED STATES

Veterinary certificate to EU

	I.1. Consignor Name	I.2. Certificate reference No I.2.a.
	Address	
		I.3. Central competent authority APHIS-VS
	Tel.	I.4. Local competent authority
	I.5. Consignee Name Address	I.6. Person responsible for the load in EU Name Address
	Postal code Tel.	Postal code Tel.
nment		
l consig	I.7. Country of ISO code I.8. Region of Code origin origin	I.9. Country of ISO I.10. Region of Code destination code destination
chec	I.11. Place of origin	I.12. Place of destination
Part I : Details of dispatched consignment	Name Approval number Address	Custom warehouse Name Approval number Address
etails		Postal code
Part I : D		
	Name Approval number Address	
	Name Approval numb er Addres s	
	I.13. Place of loading	I.14. Date of departure
	I.15. Means of transport	I.16. Entry BIP in EU
	Aeroplane	
		1.17.
	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 packages
	Ambient Chilled Frozen C	
I	Page of	

		I.2. Certificate reference No	l.2.a.
I.23. Seal/Container No		I.24. Type of packaging	
I.25. Commodities certified for:		-	
Petfood			
I.26. For transit through EU to third	d country	I.27. For import or admission into EU	
Third country	ISO code		
I.28. Identification of the commodit	ies		
Species (Scientific name)	Approval number of establishme Manufacturing plant	ents Net weight	Batch number

	11.	Health inf	ormati	on	II.a. Certificate reference No	II.b.
		l the under	sianed o	fficial veterinarian, declare	that I have read and understood Regulation (EC) No 1069/20	
	-	Parliament	and of th	e Council(1a), and in partic	ular Articles 8 and 10 thereof, and Commission Regulation (El	J) No 142/2011(^{1b}),
	II.1.	-			hapter II of Annex XIV thereto, and certify that the petfood des oved and supervised by the competent authority in accordance	
		Regulation	(EC) No	1069/2009;		
u	II.2.	nas been pi (²) either	repared (exclusively with the followir	ig animal by-products: himals slaughtered or, in the case of game, bodies or parts of	animals killed and
atio		() on the	L		consumption in accordance with Union legislation, but are not i	
Part II: Certification		(²)and/or	[-	slaughterhouse and wer	wing parts originating either from animals that have been e considered fit for slaughter for human consumption followin d the following parts of animals from game killed for hum egislation:	ng an ante-mortem
t II: C				(i) carcases or bod	ies and parts of animals which are rejected as unfit for hum Union legislation, but which did not show any signs of diseas	
ar				(ii) heads of poultry;		
Р					including trimmings and splitting thereof, horns and feet, inclu ind metacarpus bones, tarsus and metatarsus bones;	ding the phalanges
				(iv) pig bristles;		
		(²)and/or	[-	 (v) feathers;] animal by-products from 	poultry and lagomorphs slaughtered on the farm as referred	to in Article 1(3)(d)
		(),,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	L	of Regulation (EC) No 8	53/2004 of the European Parliament and of the Council (^{2a}), v nmunicable to humans or animals]	
		(²)and/or	[-	blood of animals which	did not show any signs of disease communicable through b	
					animals that have been slaughtered in a slaughterhouse tter for human consumption following an ante-mortem inspec	
		(²)and/or	[-		ng from the production of products intended for human cons s and centrifuge or separator sludge from milk processing;]	sumption, including
		(²)and/or	[-	a	n, or foodstuffs containing products of animal origin, which are	no longer intended
					for commercial reasons or due to problems of manufacturing or ich no risk to public or animal health arise;]	packaging defects
		(²)and/or	[-		ffs of animal origin, or feedingstuffs containing animal by p longer intended for feeding for commercial reasons or d	
					ging defects or other defects from which no risk to public or ani	
		(²)and/or	[-		eathers, hair, horns, hoof cuts and raw milk originating from lissease communicable through that product to humans or animations are an imparted to humans or animation of the second seco	
		(²)and/or	[-	 aquatic animals, and part communicable to human 	ts of such animals, except sea mammals, which did not show ar is or animals:1	y signs of diseases
		(²)and/or	[-		aquatic animals originating from plants or establishments man	ufacturing products
		(²)and/or	[-		iginating from animals which did not show any signs of dise	ase communicable
				(i) shells from shellf	ish with soft tissue or flesh;	
					inating from terrestrial animals:	
				-	by-products,	
				- eggs, - egg by-p	roducts, including egg shells,	
					lled for commercial reasons;]	
		(²)and/or	[-	 animal by-products from animals;] 	aquatic or terrestrial invertebrates other than species pathog	jenic to humans or
		(²)and/or		animals and parts thereo as referred to in Article 8	f of the zoological orders of Rodentia and Lagomorpha, except ι 3(a)(iii), (iv) and (v) of Regulation (EC) No 1069/2009 and Cate to (q) of that Regulation;]	
		(²)and/or	[-	material from animals w	hich have been treated with certain substances which are pro he import of the material being permitted in accordance with	
	II.3.			<u> </u>	-	
		(²)either	-	-	ent of at least 90 °C throughout its substance;]	
		(²)0r	[was (a)	in the case of animal by	dients of animal origin using exclusively products which had be /-products or derived products from meat or meat products s /C throughout its substance;	
			(b)	in the case of milk and m	C throughout its substance; nilk based products,	
			(**)		r··,	

II.	Health information		II.a. Certificate reference No	II.b.
	(i		hird countries or parts of third countries listed in column B of Ar) No 605/2010(³) submitted to a pasteurisation treatment su patase test:	
	(i	0 1 1	ed to less than 6 from third countries or parts of third countrie	es listed in column
	(of Annex I to Re	gulation (EU) No 605/2010, first submitted to a pasteurisation gative phosphatase test;	
	(i	(EU) No 605/20	hird countries or parts of third countries listed in column C of / 110, submitted to a sterilisation process or a double heat tro	
	(i	v) if they are from t	ufficient to produce a negative phosphatase test on its own; hird countries or parts of third countries listed in column C of / 10, where there has been an outbreak of foot-and-mouth dise	
			where vaccination against foot-and-mouth disease has bee inths, submitted to	n carried out in th
		either		
		- <u>a sterilis</u> or	ation process whereby an Fc value equal or greater than 3 is	achieved
		pasteuris	I heat treatment with a heating effect at least equal to sation process of at least 72 °C for at least 15 seconds and si reaction to a phosphatase test, followed by	
		either		
		heat tre	d heat treatment with a heating offect at least equal to that a atment, and which would be sufficient to produce a neg tase test, followed, in the case of dried milk, or dried milk- process	gative reaction to
		or		
		- an acidif one hou	fication process such that the pH has been maintained at lea	ss than 6 for at lea
			L,	
	si si ti	the case of gelatine, ubjected to a treatment te pH and subsequent tration and sterilisation	produced using a process that ensures that unprocessed C t with acid or alkali, followed by one or more rinses with subso t, if necessary repeated, extraction by heat, followed by puri b;	equent adjustment fication by means
	si fil (d) ir fi fi p p (i	the case of gelatine, ubjected to a treatment re pH and subsequent tration and sterilisation the case of hydrolyse inimise contamination erived from ruminant hi reparation of raw Categ) exposure of the more than 80 °C than 3,6 bar; or	produced using a process that ensures that unprocessed C t with acid or alkali, followed by one or more rinses with subset i, if necessary repeated, extraction by heat, followed by puri y; d protein produced using a production process involving app of raw Category 3 material, and, in the case of hydrolysed pro- ides and skins produced in a processing plant dedicated only i material with a molecular weight below 10000 Dalton and a p gory 3 material by brining, liming and intensive washing follow material to a pH of more than 11 for more than three hours c and subsequently by heat treatment at more than 140 °C for	equent adjustment fication by means- ropriate measures- otein entirely or par- to hydrolysed prote process involving the ved by: to at a temperature- r 30 minutes at mo
	si ## fili (d) in p p (i (i	the case of gelatine, ubjected to a treatment re pH and subsequent tration and sterilisation the case of hydrolyse inimise contamination- erived from ruminant hi roduction, using only n reparation of raw Categ exposure of the more than 80 °C than 3,6 bar; or exposure of the treatment at 140	produced using a process that ensures that unprocessed C t with acid or alkali, followed by one or more rinses with subset ; if necessary repeated, extraction by heat, followed by puri b; d protein produced using a production process involving app of raw Category 3 material, and, in the case of hydrolysed pro- ides and skins produced in a processing plant dedicated only naterial with a molecular weight below 10000 Dalton and a p gory 3 material by brining, liming and intensive washing follow material to a pH of more than 11 for more than three hours c and subsequently by heat treatment at more than 140 °C for material to a pH of 1 to 2, followed by a pH of more than 0 °C for 30 minutes at 3 bar;	equent adjustment fication by means- topriate measures tein entirely or part to hydrolysed prote process involving th ved by: at a temperature r 30 minutes at mo 11, followed by he
	si ## (d) ir # # # # (i (i (i # #	the case of gelatine, ubjected to a treatment repH and subsequent tration and sterilisation the case of hydrolyse inimise contamination arived from ruminant hi roduction, using only n reparation of raw Categ) exposure of the more than 80 °C than 3,6 bar; or) exposure of the treatment at 140 the case of egg product	produced using a process that ensures that unprocessed C t with acid or alkali, followed by one or more rinses with subso ; if necessary repeated, extraction by heat, followed by puri ; d protein produced using a production process involving app of raw Category 3 material, and, in the case of hydrolysed pro- ides and skins produced in a processing plant dedicated only naterial with a molecular weight below 10000 Dalton and a p gory 3 material by brining, liming and intensive washing follow material to a pH of more than 11 for more than three hours C and subsequently by heat treatment at more than 140 °C for material to a pH of 1 to 2, followed by a pH of more than . ⁹ °C for 30 minutes at 3 bar; ets submitted to any of the processing methods 1 to 5 or 7, as r ation (EU) No 142/2011; or treated in accordance with Chap	equent adjustment fication by means- stein entirely or part to hydrolysed prote process involving the ved by: - at a temperature- r 30 minutes at mo 11, followed by he referred to in Chapt
	si ## (ii) (d) iii # P P P (i (i) (i) (i) (e) iii # A (f) iii E (f) iii E	 the case of gelatine, ubjected to a treatment reation and sterilisation the case of hydrolyse inimise contamination erived from ruminant hi roduction, using only n reparation of raw Categ exposure of the treatment at .440 the case of egg product of Annex IV to Regulation the case of collagens a treatment involving water and the collagens a treatment involving water and the collagens 	produced using a process that ensures that unprocessed C t with acid or alkali, followed by one or more rinses with subso ; if necessary repeated, extraction by heat, followed by puri ; d protein produced using a production process involving app of raw Category 3 material, and, in the case of hydrolysed pro- ides and skins produced in a processing plant dedicated only naterial with a molecular weight below 10000 Dalton and a p gory 3 material by brining, liming and intensive washing follow material to a pH of more than 11 for more than three hours C and subsequently by heat treatment at more than 140 °C for material to a pH of 1 to 2, followed by a pH of more than . ⁹ °C for 30 minutes at 3 bar; ets submitted to any of the processing methods 1 to 5 or 7, as r ation (EU) No 142/2011; or treated in accordance with Chap	equent adjustment fication by means- ropriate measures stein entirely or par- to hydrolysed prote process involving th ved by: • at a temperature r 30 minutes at mo 11, followed by he referred to in Chapt ter II of Section X- material is subjects more rinses, filtratis
	si ti (d) in r (d) p d p p f (i (i (i (i (i (i) (i) (i) (i) (i) (i)	the case of gelatine, ubjected to a treatment e pH and subsequent tration and sterilisation the case of hydrolyse inimise contamination erived from ruminant hi roduction, using only n reparation of raw Cate) exposure of the more than 80 °C than 3,6 bar; or) exposure of the treatment at 140 the case of egg product of Annex IV to Regul of Annex IV to Regulation (the case of collagen s a treatment involving y and extrusion, the use o the case of blood prod	produced using a process that ensures that unprocessed C t with acid or alkali, followed by one or more rinses with subset ; if necessary repeated, extraction by heat, followed by puri b; d protein produced using a production process involving appro- of raw Category 3 material, and, in the case of hydrolysed pro- ides and skins produced in a processing plant dedicated only naterial with a molecular weight below 10000 Dalton and a p gory 3 material by brining, liming and intensive washing follow material to a pH of more than 11 for more than three hours c and subsequently by heat treatment at more than 140 °C for material to a pH of 1 to 2, followed by a pH of more than 0 °C for 30 minutes at 3 bar; cts submitted to any of the processing methods 1 to 5 or 7, as r ation (EU) No 142/2011; or treated in accordance with Chap (EC) No 853/2004 ; ubmitted to a process ensuring that unprocessed Category 3 washing, pH adjustment using acid or alkali followed by one or	equent adjustment fication by means- topriate measures to hydrolysed prote process involving the ved by: at a temperature r 30 minutes at mo 11, followed by he referred to in Chapt ter II of Section X- material is subjected more rinses, filtration
	(d) ir (d) ir (d) ir (d) (i (e) ir (i (e) ir (f) ir (f) ir (g) ir (g) ir (h) ir ; ;	 the case of gelatine, ubjected to a treatment e pH and subsequent tration and sterilisation the case of hydrolyse inimise contamination erived from ruminant hi roduction, using only n reparation of raw Categ exposure of the treatment at 140 exposure of the treatment at 140 the case of egg product of Annex IV to Regulation (the case of collagen s a treatment involving v at reatment involving v at reatment involving v and extrusion, the use of the case of polod provide the case of p	produced using a process that ensures that unprocessed C: t with acid or alkali, followed by one or more rinses with subset ; if necessary repeated, extraction by heat, followed by puri b; d protein produced using a production process involving appr of raw Category 3 material, and, in the case of hydrolysed pre- ides and skins produced in a processing plant dedicated only 1 naterial with a molecular weight below 10000 Dalton and a p gory 3 material by brining, liming and intensive washing follow material to a pH of more than 11 for more than three hours C and subsequently by heat treatment at more than 140 °C for material to a pH of 1 to 2, followed by a pH of more than 0 °C for 30 minutes at 3 bar; cts submitted to any of the processing methods 1 to 5 or 7, as r ation (EU) No 142/2011; or treated in accordance with Chap (EC) No 853/2004 ; submitted to a process ensuring that unprocessed Category 3 washing, pH adjustment using acid or alkali followed by one or of preservatives other than those permitted by Union legislatio ducts, produced using any of the processing methods 1 to 5 or 7.	equent adjustment fication by means- ropriate measures stein entirely or part to hydrolysed prote process involving th ved by: at a temperature r 30 minutes at mo 11, followed by he referred to in Chapt ter II of Section X- material is subject more rinses, filtration n being prohibited; or 7, as referred to ng methods 1 to 5- 5 or 7 provided th
	(d) in (d) in (d) in (f) (f) (f) (f) (f) (f) (f) (f) (f) (f)	the case of gelatine, ubjected to a treatment e pH and subsequent tration and sterilisation the case of hydrolyse inimise contamination- arived from ruminant hi roduction, using only n reparation of raw Cate() exposure of the more than 80 °C than 3.6 bar; or) exposure of the treatment at 140 the case of egg product of Annex IV to Regul on the case of collagen s a treatment involving v and extrusion, the use of the case of blood prod hapter III of Annex IV to the case of mammalia and, in the case of product and, in the case of product the case of method 7- sen applied;	produced using a process that ensures that unprocessed C t with acid or alkali, followed by one or more rinses with subset , if necessary repeated, extraction by heat, followed by puri , d protein produced using a production process involving appr of raw Category 3 material, and, in the case of hydrolysed pre- ides and skins produced in a processing plant dedicated only 1 naterial with a molecular weight below 10000 Dalton and a p gory 3 material by brining, liming and intensive washing follow material to a pH of more than 11 for more than three hours C and subsequently by heat treatment at more than 140 °C for material to a pH of 1 to 2, followed by a pH of more than 0 °C for 30 minutes at 3 bar; cts submitted to any of the processing methods 1 to 5 or 7, as r ation (EU) No 142/2011; or treated in accordance with Chap (EC) No 853/2004 ; washing, pH adjustment using acid or alkali followed by one or of preservatives other than those permitted by Union legislatio ducts, produced using any of the processing methods 1 to 5 of to Regulation (EU) No 142/2011; an processed animal protein submitted to any of the processing reine blood, submitted to any of the processing methods 1 to 5 of a heat treatment throughout its substance at a minimum temp	equent adjustment of fication by means of ropriate measures of the entirely or part to hydrolysed prote- process involving the ved by: - at a temperature of r 30 minutes at more 11, followed by here after to in Chapter the II of Section X of material is subjected more rinses, filtratic on being prohibited; for 7, as referred to ong methods 1 to 5 of 5 or 7 provided the perature of 80 °C here pomitted to any of the
	(i) (i) (i) (i) (i) (i) (i) (i)	 the case of gelatine, ubjected to a treatment e pH and subsequent tration and sterilisation the case of hydrolyse inimise contamination-arived from ruminant hi roduction, using only n reparation of raw Categ exposure of the more than 80 °C than 3,6 bar; or exposure of the treatment at 140 the case of egg product of Annex IV to Regulation (and, in the case of collagens is a treatment involving and, in the case of product the case of mammalia and, in the case of product the case of mammalia and, in the case of product the case of mammalia and, in the case of product the case of mammalia the case of mammalia the case of mammalia and, in the case of non mammalice the case of fishmeal the cas	produced using a process that ensures that unprocessed C t with acid or alkali, followed by one or more rinses with subset ; if necessary repeated, extraction by heat, followed by puri b; d protein produced using a production process involving appro- of raw Category 3 material, and, in the case of hydrolysed pro- ides and skins produced in a processing plant dedicated only inaterial with a molecular weight below 10000 Dalton and a p gory 3 material by brining, liming and intensive washing follow material to a pH of more than 11 for more than three hours c and subsequently by heat treatment at more than 140 °C for material to a pH of 1 to 2, followed by a pH of more than 0 °C for 30 minutes at 3 bar; cts submitted to any of the processing methods 1 to 5 or 7, as r ation (EU) No 142/2011; or treated in accordance with Chap (EC) No 853/2004 ; ubmitted to a process ensuring that unprocessed Category 3 washing, pH adjustment using acid or alkali followed by one or of preservatives other than those permitted by Union legislatio ducts, produced using any of the processing methods 1 to 5 or a negulation (EU) No 142/2011; an processed animal protein submitted to any of the processing methods 1 to 5 or a heat treatment throughout its substance at a minimum temp malian processed protein with the exclusion of fishmeal suf p 5 or 7 as referred to in Chapter III of Annex IV to Regulation submitted to any of the processing methods 1 to 7 as referred (EU) No 142/2011 or to a method and parameters which eni- obiological standards for derived products set out in Chapt	equent adjustment fication by means- repriate measures stein entirely or part to hydrolysed prote- process involving the ved by: - at a temperature - r 30 minutes at mo 11, followed by he referred to in Chapt ter II of Section X- material is subjected more rinses, filtration n being prohibited; or 7, as referred to - 5 or 7 provided the perature of 80 °C he omitted to any of the (EU) No 142/2011 de to in Chapter III- sure that the produ
	(d) ir (d) ir (d) ir (d) ir (e) ir (i) ir (f) ir (f	the case of gelatine, ubjected to a treatment e pH and subsequent tration and sterilisation the case of hydrolyse inimise contamination- erived from ruminant hi roduction, using only n reparation of raw Categ exposure of the more than 80 °C than 3,6 bar; or exposure of the treatment at 140 the case of egg produc of Annex IV to Regulation (the case of collagen s extreatment involving und hapter III of Annex IV to the case of collagen s extreatment involving und extrusion, the use o the case of collagen s extreatment involving und extrusion, the use o the case of mammalia and, in the case of po- the case of mammalia the case of mammalia the case of momman rocessing methods 1 to the case of fishmeal to the case of fishmeal to the case of non mam rocessing methods 1 to the case of non the case of the case of non the the case of non the the case of non the the case of non the the case of non the case of non the case of the case of non the the case of non the the case of non the case of non the case of the case of non the case of the case of the case of non the case of t	produced using a process that ensures that unprocessed C t with acid or alkali, followed by one or more rinses with subset ; if necessary repeated, extraction by heat, followed by puri b; d protein produced using a production process involving appro- of raw Category 3 material, and, in the case of hydrolysed pro- ides and skins produced in a processing plant dedicated only inaterial with a molecular weight below 10000 Dalton and a p gory 3 material by brining, liming and intensive washing follow material to a pH of more than 11 for more than three hours c and subsequently by heat treatment at more than 140 °C for material to a pH of 1 to 2, followed by a pH of more than 0 °C for 30 minutes at 3 bar; cts submitted to any of the processing methods 1 to 5 or 7, as r ation (EU) No 142/2011; or treated in accordance with Chap (EC) No 853/2004 ; ubmitted to a process ensuring that unprocessed Category 3 washing, pH adjustment using acid or alkali followed by one or of preservatives other than those permitted by Union legislatio ducts, produced using any of the processing methods 1 to 5 or a negulation (EU) No 142/2011; an processed animal protein submitted to any of the processing methods 1 to 5 or a heat treatment throughout its substance at a minimum temp malian processed protein with the exclusion of fishmeal suf p 5 or 7 as referred to in Chapter III of Annex IV to Regulation submitted to any of the processing methods 1 to 7 as referred (EU) No 142/2011 or to a method and parameters which eni- obiological standards for derived products set out in Chapt	equent adjustment fication by means- ropriate measures stein entirely or part to hydrolysed prote process involving the ved by: at a temperature- r 30 minutes at more 11, followed by he efferred to in Chapter ter II of Section X- material is subjected more rinses, filtration n being prohibited; or 7, as referred to be or 7 provided the perature of 80 °C he semitted to any of the ter II of Chapter III- sure that the produ- ter I of Annex X- hods 1 to 5 or 7 (ar in (EU) No 142/2011 (EC) No 853/200

Processed petfood other than canned petfood

 (i) ensures that all Category 3 bone material is finely crushed and degreased with hot water a treated with diute hydrochloric acid (at a minimum concentration of 4 % and a pH of less than 1 over a period of at least two days; (ii) following the procedure referred to in (i), applies a treatment of the obtained phosphoric liquor where a period of at least two days; (iii) following the procedure referred to in (i), applies a treatment of the obtained phosphoric liquor where and temperature between 30-C and 65-C; (iii) in the case of tricalcium phosphate produced by a process that ensures (i) that all Category 3 bone material is finely crushed and degreased in counter flow with hot we (bone chipe less than 14 mm); (iii) continuous cooking with steam at 145 °C during 30 minutes at 4 bar; (iii) continuous cooking with steam at 145 °C during 30 minutes at 4 bar; (iii) continuous cooking with the microbiological standards referred to in point 14.4] (iv) granulation of the tricalcium phosphate after drying in a fluid bed with air at 200 °C; (ii) in the case of flavouring inmarks, produced coording to a treatment method and parameters, which ensure that the product comples with the microbiological standards referred to in point 14.4] (²) (³) (³		Health information	i	II.a. Certificate reference No	II.b.
 treated with dilute hydrochioric acid (at a minimum concentration of 4.% and a pH of less than 1 over a period of at least two days; (ii) following the procedure referred to in (i), applies a treatment of the obtained phosphoric liquor willine, resulting in a precipitate of dicalcium phosphate at pH 4 to 7; and (iii) finally, or if drise the precipitate of dicalcium phosphate at pH 4 to 7; and (iii) finally, or if drise the precipitate of dicalcium phosphate at pH 4 to 7; and (iii) the case of tricalcium phosphate produced by a process that ensures (i) that all Category 3 bone material is finely crushed and degreased in counter-flow with hot we (bone chips less than 14 mm); (iii) continuous cooking with steam at 145 °C during 30 minutes at 4 bar; (iii) separation of the protein-broth-from the hydroxylapatite (tricalcium phosphate) by centrifugati and (iv) granulation of the tricalcium phosphate after drying in a fluid bed with air at 200 °C ; (in) in the case of flavouring innards, produced according to a treatment method and parameters, which ensure that the product complies with the microbiological standards referred to in point II.4.1 (²)or [in the case of aquatic and terrestrial invertebrates other than species pathogenic to humans or animals, has been authorised by a random sampling of at least five samples from each processed batch taken during or after storage at processing plant and complies with the following standards?): Salmonella: absence in 25; n = 5, c = 0, m = 0, M = 0, Enterobacteriaceae: n = 5, c = 2, m = 10, M = 300 in 1 gramme; thas the content is destined for feeding to pets only, bear labels indicating "NOT FOR HUMAN CONSUMPTION"; (i) the petfood is not dispatched in ready-to-sell packages on which it is clearly indica that the content is destined for between or comprise animals and does not contain and is not derived from animals be continuous/ preared an dapptere					
 (iii) finally, air dries the procipitate of dicalcium phosphate with inlet temperature of 65 °C to 325 and end temperature between 30 °C and 65 °C; (iii) in the case of tricalcium phosphate produced by a process that ensures (i) that all Category 3 bone material is finely crushed and degreased in counter flow with hot we (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 centrifugati and (iv) granulation of the tricalcium phosphate after drying in a fluid bed with air at 200 °C; (ii) in the case of flavouring innards, produced according to a treatment method and parameters, which ensithat the product complies with the microbiological standards referred to in point II.4.1] (²)or [was subject to a treatment such as drying or formentation, which has been authorised by the competent authorit (²)or [in the case of aquatic and terrestrial invertebrates other than species pathogenic to humans or animals, has be subject to a treatment which has been authorised by the competent authorit (²)or andom sampling of at least five samples from each processed batch taken during or after storage at processing plant and complies with the following standards(⁴): Salmonella: absence in 25g: n = 5, c = 0, m = 0, M = 0, Enterobacteriaceae: n = 5, c = 2, m = 10, M = 300 in 1 gramme; thas undergone all precautions to avoid contamination with pathogenic agents after treatment; was packed in new packaging, which, if the petfood is not dispatched in ready-to-sell packages on which it is clearly indica that the content is destined for feeding to pets only, bear labels indicating "NOT FOR HUMAN CONSUMPTION"; III.7. the petfood described above (²)or [is derived from other ruminants than bovine, ovine or caprine animals.] (²)or [is derived from other ruminan			treated with dilu over a period of	te hydrochloric acid (at a minimum concentration of 4 % and at least two days;	a pH of less than 1,
 (m) in the case of tricaloum phosphate produced by a process that ensures (i) that all Category 3 bone material is finely crushed and degreased in counter flow with hot we (bone chipe less than 14 mm); (ii) continuous cooking with steam at 145 °C during 30 minutes at 4 bar; (iii) continuous cooking with steam at 145 °C during 30 minutes at 4 bar; (iii) continuous cooking with steam at 145 °C during 10 minutes at 4 bar; (iii) continuous cooking with steam at 145 °C during 10 minutes at 200 °C; (iii) granulation of the tricalcium phosphate after drying in a fluid bed with air at 200 °C; (in) in the case of flavouring innards, produced according to a treatment method and parameters, which ensure that the product complies with the microbiological standards referred to in point 14.4; (²) or [in the case of aquatic and terrestrial invertebrates other than species pathogenic to humans or animals, has be subject to a treatment which has been authorised by the competent authority and which ensures that the petfood becret in a treatment					re of 65 °C to 325 °
 (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 centrifugati and (iv) granulation of the tricalcium phosphate after drying in a fluid bed with air at 200 °C.; (n) in the case of flavouring innards, produced according to a treatment method and parameters, which ensithat the product compiles with the microbiological standards: referred to in point II.4.1 (°)or [was subject to a treatment such as drying or fermentation, which has been authorised by the competent authorit (°)or [in the case of aquatic and terrestrial invertebrates other than species pathogenic to humans or animals, has be subject to a treatment which has been authority and which ensures that the petfor poses no unacceptable risks to public and animal health.] 4. was analysed by a random sampling of at least five samples from each processed batch taken during or after storage at processing plant and complies with the following standards(°): Salmonella: absence in 25g: n = 5, c = 0, m = 0, M = 0, Enterobacteriaceae: n = 5, c = 2, m = 10, M = 300 in 1 gramme; 5. has undergone all precautions to avoid contamination with pathogenic agents after treatment; 6. was packed in new packaging, which, if the petfood is not dispatched in ready-to-sell packages on which it is clearly indicat that the content is destined for feeding to pets only, bear labels indicating "NOT FOR HUMAN CONSUMPTION"; [II.7. the petfood described above (°) either [bovine, ovine or caprine animals.] (°)or [is derived from other ruminants than bovine, ovine or caprine animals.] (°)or [is derived from other ruminants than bovine, ovine or coprine animals.] (°)or [is derived from other ruminants than bovine.] (°)or [is derived from other ruminants than bovine.] (°)or [is derived from other ruminants than bovine.]<td></td><td>(m)</td><td>•</td><td></td><td></td>		(m)	•		
 (iii) separation of the protein broth from the hydroxyapatite (tricalcium phosphate) by centrifugatiand (iv) granulation of the tricalcium phosphate after drying in a fluid bed with air at 200 °C.; (n) in the case of flavouring innards, produced according to a treatment method and parameters, which ensithat the product complies with the microbiological standards referred to in point II.4.] (²)or [was subject to a treatment such as drying or formentation, which has been authorised by the competent authorif (³)or [in the case of aquatic and terrestrial invertebrates other than species pathogenic to humans or animals, has be subject to a treatment which has been authorised by the competent authorif (⁴)or [in the case of aquatic and terrestrial invertebrates other than species pathogenic to humans or animals, has be subject to a treatment which has been authorised by the competent authorif (⁴)or [in the case of aquatic and terrestrial invertebrates other than species pathogenic to humans or animals, has be subject to a treatment which has been authorised by the competent authorif was analysed by a random sampling of at least five samples from each processed batch taken during or after storage at processing plant and complies with the following standards(⁴): Salmonella: absence in 25g: n = 5, c = 0, m = 0, M = 0, Enterobacteriaceae: n = 5, c = 2, m = 10, M = 300 in 1 gramme; has undergone all precautions to avoid contamination with pathogenic agents after treatment; was packed in new packaging, which, if the petfood is not dispatched in ready-to-sell packages on which it is clearly indicat that the content is destined for feeding to pets only, bear labels indicating "NOT FOR HUMAN CONSUMPTION"; (III.7. the petfood described above (²)either [is derived from other ruminants than bovine, ovine or caprine animals.] (²)or [is derived from other ruminants than bovine, ovine or caprine animals.] (²)or [is derived from		•			er-flow with hot wate
 (iv)granulation of the tricalcium phosphate after drying in a fluid bed with air at 200 °C.; (n)			(iii) separation of th	.	ate) by centrifugation
 that the product complies with the microbiological standards referred to in point II.4.] (²)or [was subject to a treatment such as drying or fermentation, which has been authorised by the competent authori (²)or [in the case of aquatic and terrestrial invertebrates other than species pathogenic to humans or animals, has be subject to a treatment which has been authorised by the competent authority and which ensures that the petfor poses no unacceptable risks to public and animal health.] 4. was analysed by a random sampling of at least five samples from each processed batch taken during or after storage at processing plant and complies with the following standards^(*): Salmonella: absence in 25g: n = 5, c = 0, m = 0, M = 0, Enterobacteriaceae: n = 5, c = 2, m = 10, M = 300 in 1 gramme; 5. has undergone all precautions to avoid contamination with pathogenic agents after treatment; 6. was packed in new packaging, which, if the petfood is not dispatched in ready-to-sell packages on which it is clearly indicat that the content is destined for feeding to pets only, bear labels indicating "NOT FOR HUMAN CONSUMPTION"; (III.7. the petfood described above (²)either [is derived from other ruminante than bovine, ovine or caprine animals.] (²)or [is derived from other ruminante than bovine, ovine or caprine animals.] (²)or [is derived from bovine, ovine and caprine materials other than those derived from: (²) either [bovine, ovine and caprine materials other than those derived from animals be continuously reared and slaughtered in a country or region classified as posing a negligit BSE risk in accordance with Decision 2007/453/EC.]] (²)or [(a) specified risk material as defined in point 1 of Annex V to Regulation (EC) 999/2001 of the European Parliament and of the Council(⁶); (b) mechanically separated meat obtained from bones of bovine, ovine or capr animals, except from those animals that were born, continu				e tricalcium phosphate after drying in a fluid bed with air at 2	200 °C ;
 (²)or [in the case of aquatic and terrestrial invertebrates other than species pathogenic to humans or animals, has be subject to a treatment which has been authorised by the competent authority and which ensures that the petfor poses no unacceptable risks to public and animal health.] 4. was analysed by a random sampling of at least five samples from each processed batch taken during or after storage at processing plant and complies with the following standards(⁴): Salmonella: absence in 25g: n = 5, c = 0, m = 0, M = 0, Enterobacteriaceae: n = 5, c = 2, m = 10, M = 300 in 1 gramme; 5. has undergone all precautions to avoid contamination with pathogenic agents after treatment; 6. was packed in new packaging, which, if the petfood is not dispatched in ready-to-sell packages on which it is clearly indicat that the content is destined for feeding to pets only, bear labels indicating "NOT FOR HUMAN CONSUMPTION"; [II.7. the petfood described above (²)or [is derived from other ruminants than bovine, ovine or caprine animals.] (²)or [is derived from bovine, ovine or caprine animals and does not contain and is not derived from: (³) either [bovine, ovine and caprine materials other than those derived from animals be continuously reared and slaughtered in a country or region classified as posing a negligi BSE-risk in accordance with Decision 2007/453/EC.] (²)or [(a) specified risk material as defined in point 1 of Annex V to Regulation (EC) 999/2001 of the European Parliament and of the Council(⁵); (b) mechanically separated meat obtained from bones of bovine, ovine or caprine animals, except from those animals that were born, continuously reared a slaughtered in a country or region classified as posing a negligible BSE risk accordance with Commission Decision 2007/453/EC(⁶), in which there has be no indigenous BSE case, 					
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 5. has undergone all precautions to avoid contamination with pathogenic agents after treatment; 6. was packed in new packaging, which, if the petfood is not dispatched in ready-to-sell packages on which it is clearly indicat that the content is destined for feeding to pets only, bear labels indicating "NOT FOR HUMAN CONSUMPTION";)[II.7. the petfood described above (²)<i>either</i> (is derived from other ruminants than bovine, ovine or caprine animals.] (²)<i>or</i> (is derived from bovine, ovine or caprine animals and does not contain and is not derived from: (²)<i>either</i> (²)<i>either</i> (²)<i>either</i> (²)<i>either</i> (²)<i>either</i> (²)<i>either</i> (²)<i>either</i> (²)<i>or</i> (³) (²)<i>or</i> (⁴) (²)<i>or</i> (⁴) (²)<i>or</i> (⁴) (²)<i>or</i> (¹) (²)<i>or</i> (¹) (²) (³) (⁴) (0		
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		()01	. ,	999/2001 of the European Parliament and of the Council ⁽⁵); mechanically separated meat obtained from bones of bov	c ()
animals which have been killed, after stunning, by laceration of the central nervo tissue by means of an elongated rod-shaped instrument introduced into the crar cavity, or by means of gas injected into the cranial cavity, except for those anim that were born, continuously reared and slaughtered in a country or reg				slaughtered in a country or region classified as posing a r	negligible BSE risk
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Page ____ of ____

II.	Health information	II.a. C	ertificate reference N	NU	II.b.
	 Box reference I.28: Species: select from Pesca, Mollusca, Crustacea, Invertebra 				r than Ruminantia or Suid
Part I					
(1a)	OJ L 300, 14.11.2009, p. 1.				
(1b)	OJ L 54, 26.2.2011, p. 1.				
(2)	Delete as appropriate.				
(^{2a})	OJ L 139, 30.4.2004, p. 55.				
(^{2b})	OJ L 125, 23.5.1996, p. 3.				
(3)	OJ L 175, 10.7.2010, p. 1.				
(4)	Where:				
	n = number of samples to be tested;				
	m = threshold value for the number of does not exceed m;			-	
	M = maximum value for the number of more samples is M or more; and			-	
(E)	c = number of samples the bacterial of if the bacterial count of the other s			۸, the sample still b،	eing considered accept
(5)	OJ L 147, 31.5.2001, p. 1.				
(6)	OJ L 172, 30.6.2007, p. 84.				
	 The signature and the stamp must be in the person responsible for the person responsible				anly for votoring a surra
	 Note for the person responsible for the and must accompany the consignment 	0			
Offic	ial veterinarian/Official inspector				
2.110					
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	Name (in capital letters):	Qualif	ication and title:		
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