



Turkije, petfood

Code: **DPDL-154** Versie: 1.1.6

Ingangsdatum: 15-11-2023

Eigenaar: NVWA O&O, team Export

Versie	Datum	Wijziging ten opzichte van vorige versie
1.1.4	13-12-2022	Door de autoriteiten van Turkije is aangegeven dat voor export van smaakgevende ingewanden het certificaat voor processed petfood other than canned petfood moet worden gebruikt.
1.1.5	01-05-2023	In het certificaat voor verwerkt petfood anders dan ingeblikt moeten in verklaring II.3, optie 2, de niet van toepassing zijnde deelverklaringen worden doorgehaald. In de certificaten in veld I.8 worden vermeld "regionalisation not applicable".
1.1.6	15-11-2023	In de certificaten in veld I.8 worden vermeld de ISO-code van het land. Aanpassing veld I.25 naar Petfood.

1 Doel en toepassingsgebied

Deze instructie geldt voor het exporteren van petfood naar Turkije.

De instructie beschrijft de voorwaarden die gelden voor de invoer in Turkije, de controles die de NVWA hiervoor moet uitvoeren, en de gegevens die het bedrijfsleven moet aanleveren aan de NVWA. Over de certificeringseisen die gelden voor de export van petfood naar Turkije zijn officiële bilaterale afspraken gemaakt. Deze afspraken zijn bindend, van deze afspraken kan dus niet worden afgeweken.

2 Wettelijke basis

2.1 EU-regelgeving

- Verordening (EG) nr. 183/2005
- Verordening (EG) nr. 1069/2009
- Verordening (EU) nr. 142/2011

2.2 Nationale wetgeving

- Wet dieren
- Besluit dierlijke producten
- Regeling dierlijke producten

Overige

- Bilaterale afspraken tussen Turkije en Nederland.

3 Definities

n.v.t.

4 Werkwijze

De export van petfood naar Turkije is toegestaan.

De autoriteiten in Turkije eisen dat de geëxporteerde partijen petfood te koppelen zijn aan de exportcertificaten. Op basis hiervan eisen de Turkse autoriteiten dat bij verzending per schip containernummer en zegelnummer worden vermeld op het certificaat.

Voor zendingen die per vrachtwagen naar Turkije gaan is het vermelden van kenteken en zegelnummer op de exportcertificaten verplicht.

Zonder vermelding van de betreffende gegevens kan een exportcertificaat niet afgegeven worden.

In april 2014 is door Turkije aangegeven dat bovenstaande geldt voor:

- petfood dat vleesgrondstoffen bevat (in iedere vorm en in ieder percentage),
- petfood met meer dan 50% zuivel, en
- plantaardige producten, geëxporteerd met een fytosanitair certificaat.

Deze eisen gelden dus niet langer voor plantaardige producten die met een veterinair certificaat worden geëxporteerd.

Turkije heeft aangegeven voor afgegeven certificaten de volgende geldigheidstermijnen te hanteren:

- Voor zendingen per vliegtuig 7 dagen vanaf afgiftedatum;
- Voor zendingen over de weg 15 dagen vanaf afgiftedatum;
- Voor zendingen per trein 30 dagen vanaf afgiftedatum;
- Voor zeevracht 60 dagen vanaf afgiftedatum.

4.1 Canned petfood

Certificaat: zie *bijlage 1*.

Algemeen:

- Raadpleeg vooraf de instructie Tijdelijke Maatregelen Derde Landen (TMDL-01) op mogelijke exportbeperkingen. Als in de TMDL-01 informatie staat die in strijd is met een landeninstructie dan is de informatie vermeld in de TMDL-01 leidend.
- Indien belanghebbende de gevraagde garanties niet kan aanleveren, zal de certificering niet doorgaan.

Invuleisen:

- I.8.: Dit veld hoeft alleen te worden ingevuld indien er sprake is van regionalisatie; zolang dit niet aan de orde is wordt er standaard de ISO-code van het land ingevuld.
- I.11.: In het certificaat kan slechts één producent worden opgenomen. Alleen producten van de betreffende producent kunnen op één certificaat gecertificeerd worden.
- I.19.: De HS-code voor ingeblikt petfood is 2309; dit moet in e-CertNL op orderniveau worden ingevoerd.
- I.28.: In geval er meerdere soorten dierlijke grondstoffen in het product zijn verwerkt, dan moet de diersoort op grondstofniveau ingevuld worden, om te kunnen tonen op het certificaat.

Toelichting bij het certificaat:

Aanhef:

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 and in particular Article 8 and 10 thereof, and Commission Regulation (EU) No 142/2011, and in particular Chapter II of Annex XIII and Chapter II of Annex XIV thereof and certify that the petfood described above:

Deze verklaring kan worden afgegeven indien de certificerende NVWA-dierenarts kennis heeft genomen van de relevante bepalingen van de genoemde EU-regelgeving (Verordening (EG) nr. 1069/2009 en Verordening (EG) nr. 142/2011).

Verklaring II.1:

has been prepared and stored in an establishment or plant approved and supervised by the competent authority in accordance with Article 24 of Regulation (EC) No 1069/2009;

Deze verklaring kan worden afgegeven voor een product vervaardigd in een bedrijf in Nederland of in een andere EU-lidstaat met een erkenning op basis van Verordening (EG) nr. 1069/2009, of vervaardigd in een bedrijf in een derde land en via een zichtkeuring binnen de EU gebracht en

verzonden vanaf een bedrijf met een registratie of een erkenning op basis van Verordening (EG) nr. 183/2005.

Verklaring II.2:

has been prepared exclusively with the following animal by-products:

- either⁽¹⁾ carcasses and parts of animal slaughtered or, in the case of game, bodies or parts of animal killed, and which are fit for human consumption, but are not intended for human consumption for commercial reasons;*
- and/or⁽¹⁾ carcasses 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:*
- (i) carcasses or bodies and parts of animals which are rejected as unfit for human consumption, 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;*
- and/or⁽¹⁾ animal by products from poultry and lagomorphs slaughtered on the farm as referred to in Article 1(3)(d) of Regulation (EC) No 853/2004, which did not show any signs of disease communicable to humans or animals;*
- and/or⁽¹⁾ blood of animals which did not show any signs of disease communicable through blood to humans or animals, obtained from animals that have been slaughtered in a slaughterhouse after having been considered fit for slaughter for human consumption following an ante-mortem inspection;*
- and/or⁽¹⁾ animal by-products arising from the production of products intended for human consumption, including degreased bone, greaves and centrifuge or separator sludge from milk processing;*
- and/or⁽¹⁾ products of animal origin or foodstuffs containing products of animal origin, which are no longer intended for human consumption for commercial reasons or due to problems of manufacturing or packaging defects or other defects from which no risk to public or animal health arise;*
- and/or⁽¹⁾ petfood and feedingstuffs of animal origin, or feedingstuffs containing animal by-products or derived products, which are no longer intended for feeding for commercial reasons or due to problems of manufacturing or packaging defects or other defects from which no risk to public or animal health arises;*
- and/or⁽¹⁾ blood, placenta, wool, feathers, hair, horns, hoof cuts and raw milk originating from live animals that did not show signs of any disease communicable through that product to humans or animals;*
- and/or⁽¹⁾ aquatic animals, and parts of such animals, except sea mammals, which did not show any signs of diseases communicable to humans or animals;*
- and/or⁽¹⁾ animal by-products from aquatic animals originating from plants or establishments manufacturing products for human consumption;*
- and/or⁽¹⁾ 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 following originating from terrestrial animals;*
 - hatchery by-products;*
 - eggs;*
 - egg by-products, including egg shells;*
 - (iii) day-old chicks killed for commercial reasons;*
- and/or⁽¹⁾ animal by-products from aquatic or terrestrial invertebrates other than species pathogenic to humans or animals;*
- and/or⁽¹⁾ animals and parts thereof of the zoological orders of Rodentia and Lagomorpha, except Category 1 material as referred to in Article 8(a)(iii), (iv) and (v) of Regulation (EC) No 1069/2009 and Category 2 material as referred to in Article 9(a) to (g) of that Regulation;*

and/or⁽¹⁾ material from animals which have been treated with certain substances which are prohibited pursuant to Directive 96/22/EC, the import of the material being permitted in accordance with Article 35(a)(ii) of Regulation (EC) No 1069/2009;

Deze verklaring kan worden afgegeven op basis van een grondstoffenlijst. Belanghebbende moet de herkomst van de grondstoffen en de aard van het cat. 3-materiaal aantonen. De niet van toepassing zijnde opties moeten worden doorgehaald.

De aanvrager dient zelf in de aanvraag in e-CertNL onder het tabblad "documenten" bij "verklaringsteksten" de keuze te maken welke tekst getoond wordt.

Voor een product vervaardigd in Nederland of in een andere EU-lidstaat kan deze verklaring worden afgegeven op basis van een grondstoffenlijst opgesteld door het productiebedrijf.

Voor een product vervaardigd in een derde land kan deze verklaring worden afgegeven door het overleggen van het EU-importcertificaat met daarin een verklaring van gelijke strekking plus bijbehorend GGB.

Verklaring II.3:

has been subjected to heat treatment to a minimum F_c value of 3 in hermetically sealed containers;

Deze verklaring kan worden afgegeven op basis van EU- en nationale regelgeving voor ingeblikt petfood, afkomstig van een productiebedrijf met een erkenning op basis van Verordening (EG) nr. 1069/2009.

Verklaring II.4:

was analysed by a random sampling of at least five samples from each processed batch by laboratory diagnostic methods to ensure adequate heat treatment of the whole consignment as foreseen under point II.3;

Deze verklaring kan worden afgegeven op basis van een bombagetest van vijf blikjes/gesloten recipiënten uit de te exporteren partij met negatieve uitslag, aan te leveren door belanghebbende. Uit de laboratoriumuitslag moet middels een partij- of batchnummer blijken dat de uitslag gerelateerd is aan de te exporteren partij. De uitslag mag maximaal vier weken oud zijn.

Voor een product vervaardigd in een andere EU-lidstaat kan deze verklaring worden afgegeven op basis van een veterinaire verklaring van gelijke strekking of door het overleggen van uitslagen van microbiologisch onderzoek op basis van partijbemonstering in het Nederlandse opslagbedrijf.

Voor een product vervaardigd in een derde land kan deze verklaring worden afgegeven door het overleggen van het EU-importcertificaat met daarin verklaringen van gelijke strekking plus bijbehorend GGB of door het overleggen van uitslagen van microbiologisch onderzoek op basis van partijbemonstering in het Nederlandse opslagbedrijf.

Verklaring II.5:

has undergone all precautions to avoid contamination with pathologic agents after treatment;

Deze verklaring kan worden afgegeven op basis van EU- en nationale regelgeving voor een product vervaardigd in een bedrijf met een registratie of erkenning op basis van Verordening (EG) nr. 1069/2009 of afkomstig van een bedrijf met een registratie op basis van Verordening (EG) nr. 183/2005.

Verklaring II.6:

The petfood described above⁽¹⁾

either⁽¹⁾ is derived from other ruminants than bovine, ovine or caprine animals;

or⁽¹⁾ is derived from bovine, ovine or caprine animals and does not contain and is not derived from:

either⁽¹⁾ bovine, ovine or caprine materials other than those derived from animals born, continuously reared and slaughtered in a country or region which is classified as a "negligible" risk status by WOAH;

or⁽¹⁾ (a) specified risk material as defined in point 1 of Annex V to Regulation (EC) No 999/2001 of the European Parliament and of the Council;

(b) mechanically separated meat obtained from bones of bovine, ovine or caprine animals, except from those animals that were born, continuously reared and slaughtered in a country or region classified as posing a negligible BSE risk in accordance with WOAH, in which there has been no 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 WOAH.

Deze verklaring kan worden afgegeven na controle. De niet-relevante deelverklaringen moeten worden doorgehaald. Belanghebbende moet aantonen van welke grondstoffen het ingeblikt petfood is gemaakt (diersoort en herkomst). Deze verklaring is alleen van toepassing voor herkauwergondstoffen verkregen van het slachtproces en niet voor andere herkauwergondstoffen zoals zuivelgrondstoffen. Indien geen andere herkauwergondstoffen dan afkomstig van zuivel zijn verwerkt, moet de gehele verklaring worden doorgehaald.

Voor een product vervaardigd in Nederland of in een andere EU-lidstaat kan betreffende deelverklaring worden afgegeven op basis van een verklaring van gelijke strekking opgesteld door het productiebedrijf.

Voor een product vervaardigd in een derde land kan betreffende deelverklaring worden afgegeven door het overleggen van het EU-importcertificaat met daarin een verklaring van gelijke strekking plus bijbehorend GGB.

Voor een product vervaardigd in Nederland of in een andere EU-lidstaat geldt dat de items (a), (b) en (c) van de tweede deelverklaring kunnen worden afgegeven op basis van EU- en nationale regelgeving.

4.2 Other than canned petfood, inclusief smaakgevende ingewanden

Certificaat: zie *bijlage 2*.

Algemeen:

- Raadpleeg vooraf de instructie Tijdelijke Maatregelen Derde Landen (TMDL-01) op mogelijke exportbeperkingen. Als in de TMDL-01 informatie staat die in strijd is met een landeninstructie dan is de informatie vermeld in de TMDL-01 leidend.
- Door de autoriteiten van Turkije is aangegeven dat voor export van smaakgevende ingewanden het certificaat voor processed petfood other than canned petfood moet worden aangevraagd.
- Indien belanghebbende de gevraagde garanties niet kan aanleveren, zal de certificering niet doorgaan.

Invuleisen:

- I.8.: Dit veld hoeft alleen te worden ingevuld indien er sprake is van regionalisatie; zolang dit niet aan de orde is wordt er standaard de ISO-code van het land ingevuld.
- I.11.: In het certificaat kan slechts één producent worden opgenomen. Alleen producten van de betreffende producent kunnen op één certificaat gecertificeerd worden.
- I.19.: De HS-code voor niet ingeblikt petfood is 2309; Alleen petfood wat onder deze code valt mag worden gecertificeerd. (Deze opmerking naar aanleiding van de voetnoot waarin een aantal HS-codes voor producten/grondstoffen is opgevoerd, welke zouden kunnen worden geëxporteerd. Aangezien het certificaat bestemd is voor petfood (als eindproduct), zijn deze overige HS-codes niet van toepassing.)
- I.28.: In geval er meerdere soorten dierlijke grondstoffen in het product zijn verwerkt, dan moet de diersoort op grondstofniveau ingevuld worden, om te kunnen tonen op het certificaat.

Toelichting bij het certificaat:

Aanhef:

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 and in particular Articles 8 and 10 thereof, and Commission Regulation (EU) No 142/2011, and in particular Chapter II of Annex XIII and Chapter II of Annex XIV thereto and certify that the petfood described above:

Deze verklaring kan worden afgegeven indien de certificerende NVWA-dierenarts kennis heeft genomen van de relevante bepalingen van de genoemde EU-regelgeving (Verordening (EG) nr. 1069/2009 en Verordening (EG) nr. 142/2011).

Verklaring 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;

Deze verklaring kan worden afgegeven voor een product vervaardigd in een bedrijf in Nederland of in een andere EU-lidstaat met een erkenning op basis van Verordening (EG) nr. 1069/2009, of vervaardigd in een bedrijf in een derde land en via een zichtkeuring binnen de EU gebracht en verzonden vanaf een bedrijf met een registratie of een erkenning op basis van Verordening (EG) nr. 183/2005.

Verklaring II.2:

Has been prepared exclusively with the following animal by-products:

- either⁽¹⁾ Carcasses and parts of animal slaughtered or, in the case of game, bodies or parts of animal killed, and which are fit for human consumption, but are not intended for human consumption for commercial reasons;*
- and/or⁽¹⁾ Carcasses 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:*
 - i) carcasses or bodies and parts of animals which are rejected as unfit for human consumption, 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;*
- and/or⁽¹⁾ Animal by-products from poultry and lagomorphs slaughtered on the farm as referred to in Article 1(3)(d) of Regulation (EC) No 853/2004, which did not show any signs of disease communicable to humans or animals;*
- and/or⁽¹⁾ Blood of animals which did not show any signs of disease communicable through blood to humans or animals, obtained from animals that have been slaughtered in a slaughterhouse after having been considered fit for slaughter for human consumption following an ante-mortem inspection;*
- and/or⁽¹⁾ Animal by-products arising from the production of products intended for human consumption, including degreased bone, greaves and centrifuge or separator sludge from milk processing;*
- and/or⁽¹⁾ Petfood and feedingstuffs of animal origin, or feedingstuffs containing animal by-products or derived products, which are no longer intended for feeding for commercial reasons or due to problems of manufacturing or packaging defects or other defects from which no risk to public or animal health arises;*
- and/or⁽¹⁾ Blood, placenta, wool, feathers, hair, horns, hoof cuts and raw milk originating from live animals that did not show signs of any disease communicable through that product to humans or animals;*
- and/or⁽¹⁾ Aquatic animals, and parts of such animals, except sea mammals, which did not show any signs of diseases communicable to humans or animals;*
- and/or⁽¹⁾ Animal by-products from aquatic animals originating from plants or establishments manufacturing products for human consumption;*
- and/or⁽¹⁾ 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 following originating from terrestrial animals:
 - hatchery by-products;
 - eggs;
 - egg by-products, including egg shells;*
 - iii) day-old chicks killed for commercial reasons;*
- and/or⁽¹⁾ Animal by-products from aquatic or terrestrial invertebrates other than species pathogenic to humans or animals;*
- and/or⁽¹⁾ Animals and parts thereof of the zoological orders of Rodentia and Lagomorpha, except Category 1 material as referred to in Article 8(a)(iii), (iv) and (v) of Regulation (EC) No 1069/2009 and Category 2 material as referred to in Article 9(a) to (g) of that Regulation;*

and/or⁽¹⁾ Material from animals which have been treated with certain substances which are prohibited pursuant to Directive 96/22/EC, the import of the material being permitted in accordance with Article 35(a)(ii) of Regulation (EC) No 1069/2009;

Deze verklaring kan worden afgegeven op basis van een grondstoffenlijst. Belanghebbende moet de herkomst van de grondstoffen en de aard van het categorie 3-materiaal aantonen. De niet van toepassing zijnde opties kunnen worden doorgehaald.

De aanvrager dient zelf in de aanvraag in e-CertNL onder het tabblad "Documenten" bij "Verklaringsteksten" de keuze te maken welke tekst getoond wordt.

Voor een product vervaardigd in Nederland of in een andere EU-lidstaat kan deze verklaring worden afgegeven op basis van een grondstoffenlijst opgesteld door het productiebedrijf.

Voor een product vervaardigd in een derde land kan deze verklaring worden afgegeven door het overleggen van het EU-importcertificaat met daarin een verklaring van gelijke strekking plus bijbehorend GGB.

Verklaring II.3:

- either⁽¹⁾ Was subjected to a heat treatment of at least 90°C throughout its substance;*
- or⁽¹⁾ Was produced as regards ingredients of animal origin using exclusively products which had been:*
 - a. In the case of animal by-products or derived products from meat or meat products subjected to a heat treatment of at least 90°C throughout its substance;*
 - b. In the case of milk and milk based products,*
 - i) Coming from country/region where are free from foot-and-mouth disease; and raw milk used in the product was treated with a heating effect at least equivalent to that achieved by a pasteurisation process of at least 72°C for 15 seconds to produce a negative phosphatase test;*
 - ii) Coming from a country/region where has not been occurred any foot-and-mouth disease for last 12 months and pH of raw milk used in the product was reduced to less than 6 and firstly submitted to a pasteurisation treatment sufficient to produce a negative phosphatase test;*
 - iii) Coming from a country/region where has not been occurred any foot-and-mouth disease for last 12 months and raw milk used in the product was submitted to a sterilisation process or a double heat treatment where each treatment was sufficient to produce a negative phosphatase test*
 - iv) Coming from a country where has been occurred foot-and-mouth disease for last 12 months or where vaccination against foot-and-mouth disease has been carried out in the last 12 months submitted to:*
 - either A sterilisation process whereby an F_c value equal or greater than 3 is achieved;*
 - or An initial heat treatment with a heating effect at least equal to that achieved by a pasteurisation process of at least 72°C for at least 15 seconds and sufficient to produce a negative reaction to a phosphatase test, followed by:*
 - either: A second heat treatment with a heating effect at least equal to that achieved by the initial heat treatment, and which would be sufficient to produce a negative reaction to a phosphatase test, followed, in the case of dried milk, or dried milk-based products by a drying process;*
 - or An acidification process such that the pH has been maintained at less than 6 for at least one hour;*
 - c. In the case of gelatine, produced using a process that ensures that unprocessed Category 3 material is subjected to a treatment with acid or alkali, followed by one or more rinses with subsequent adjustment of the pH and subsequent, if necessary repeated, extraction by heat, followed by purification by means of filtration and sterilisation;*
 - 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 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;
 - e. In the case of egg products submitted to any of the processing methods 1 to 5 or 7 as referred to in Chapter III of Annex IV to Regulation (EU) No 142/2011 or treated in accordance with Chapter II of Section X of Annex III to Regulation (EC) No 853/2004 of the European Parliament and of the Council;
 - f. In the case of collagen submitted to a process ensuring that unprocessed Category 3 material is subjected to a treatment involving washing, pH adjustment using acid or alkali followed by one or more rinses, filtration and extrusion, the use of preservatives other than those permitted by Union legislation being prohibited;
 - g. In the case of blood products produced using any of the processing methods 1 to 5 or 7, as referred to in Chapter III of Annex IV to Regulation (EU) No 142/2011;
 - h. In the case of mammalian processed animal protein submitted to any of the processing methods 1 to 5 or 7 and in the case of porcine blood, submitted to any of the processing methods 1 to 5 or 7 provided that in the case of method 7 a heat treatment throughout its substance at a minimum temperature of 80°C has been applied;
 - i. In the case of non-mammalian processed protein with the exclusion of fishmeal submitted to any of the processing methods 1 to 5 or 7 as referred to in Chapter III of Annex IV to Regulation (EU) No 142/2011;
 - j. In the case of fishmeal submitted to any of the processing methods 1 to 7 as referred to in Chapter III of Annex IV to Regulation (EU) No 142/2011 or to a method and parameters which ensure that the products complies with the microbiological standards for derived products set out in Chapter I of Annex X to Regulation (EU) No 142/2011;
 - k. In the case of rendered fat including fish oils, submitted to any of the processing methods 1 to 5 or 7 (and method 6 in the case of fish oil) as referred to in Chapter III of Annex IV to Regulation (EU) No 142/2011 or produced in accordance with Chapter II of Section XII of Annex III to Regulation (EC) No 853/2004; rendered fats from ruminant animals must be purified in such a way that the maximum level of remaining total insoluble impurities does not exceed 0,15% in weight;
 - l. In the case 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;
 - and ii) Following the procedure under (i), applied a treatment of the obtained phosphoric liquor with lime, resulting in a precipitate of dicalcium phosphate at pH 4 to 7;
 - 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 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 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. In the case of flavouring innards, produced according to treatment method and parameters, which ensure that the product complies with the microbiological standards referred to under point II.4;
- or⁽¹⁾ Was subject to a treatment such as drying or fermentation, which has been authorised by the competent authority;
- or⁽¹⁾ In the case of aquatic and terrestrial invertebrates other species pathogenic to humans or animals, be subject to a treatment which has been authorised by the competent authority and ensures that the pet food poses no unacceptable risks to public and animal health;
- Deze verklaring kan worden afgegeven na controle. Belanghebbende moet aantonen dat het product de genoemde hittebehandeling(en) heeft ondergaan door middel van gegevens met betrekking tot het productieproces.
- Er zijn vier opties mogelijk; bij optie 2 gelden verschillende procesparameters voor de verschillende soorten grondstoffen.

- Optie 1 is van toepassing in geval het product zelf 90 °C in de kern is verhit.
- Optie 2 is van toepassing in geval het product niet 90 °C in de kern is verhit, maar de verschillende grondstoffen van dierlijke origine wel met een toegestane behandelmethode zijn verkregen. Bij optie 2 gelden verschillende procesparameters voor de verschillende soorten grondstoffen; alleen voor de gebruikte grondstoffen van dierlijke origine moet worden aangetoond dat de genoemde behandelmethode is toegepast.
De niet van toepassing zijnde sub-opties moeten worden doorgehaald.
Voor deelverklaring a en c t/m n dient de aanvrager zelf in de aanvraag in e-CertNL onder het tabblad "Documenten" bij "Verklaringsteksten" de keuze te maken welke tekst getoond wordt.
Wanneer zuivelgrondstoffen zijn gebruikt (deelverklaring b) moet in de aanvraag worden verklaard welke optie van toepassing is op basis van de toegepaste hittebehandeling en de MKZ status van het herkomstland van de zuivelgrondstof.
- Optie 3 is van toepassing in geval het product is behandeld met een door de autoriteit goedgekeurde andere methode, zoals bijvoorbeeld drogen of fermentatie.
- Optie 4 is van toepassing voor aquatische of terrestrische ongewervelde dieren in geval deze zijn behandeld met een door de autoriteit goedgekeurde methode.

Het is niet mogelijk om op één certificaat producten waarbij verschillende opties van toepassing zijn te exporteren.

Voor een product vervaardigd in een productiebedrijf in Nederland kan de behandelmethode worden onderbouwd op basis van een bedrijfsverklaring met betrekking tot het productieproces **afkomstig van de producent van het product**, hetgeen periodiek wordt geverifieerd.

Voor een product vervaardigd in een andere EU-lidstaat moet de behandelmethode als volgt worden onderbouwd.

- Wanneer gekozen wordt voor optie 1 of 2 (waarbij in geval van optie 2 geen sprake is van zuivelgrondstoffen), kan dit op basis van een bedrijfsverklaring van gelijke strekking opgesteld door de producent.
- Wanneer gekozen wordt voor optie 2 en zuivelgrondstoffen zijn gebruikt, moet dit op basis van een veterinaire verklaring van gelijke strekking opgesteld door de bevoegde autoriteit. Dit kan zijn in de vorm van een certificaat bij iedere ontvangen zending, dan wel een 'long term declaration' (maximaal twaalf maanden oud).
- Wanneer gekozen wordt voor optie 3 of 4 moet dit op basis van een veterinaire verklaring van gelijke strekking opgesteld door de bevoegde autoriteit. Dit kan zijn in de vorm van een certificaat bij iedere ontvangen zending, dan wel een 'long term declaration' (maximaal twaalf maanden oud).

Voor een product vervaardigd in een derde land moet de behandelmethode worden onderbouwd door het overleggen van het EU-importcertificaat met daarin een verklaring van gelijke strekking plus bijbehorend GGB.

Verklaring II.4:

Were analysed by a random sampling of at least five samples from each processed batch taken during or after storage at the processing plant and complies with the following standards(2):

- *Salmonella: absence in 25 g: n = 5, c = 0, m = 0, M = 0*
- *Enterobacteriaceae: n = 5, c = 2, m = 10, M=300 in 1 gram;*

Deze verklaring kan worden afgegeven op basis van laboratoriumuitslagen van vijf deelmonsters voor de genoemde pathogenen, aan te leveren door belanghebbende. Dit mag als volgt worden geïnterpreteerd: per certificaataanvraag moet één product per productiebedrijf zijn onderzocht (vijf deelmonsters, waarbij Salmonella door het laboratorium als mengmonster mag worden onderzocht). Uit de laboratoriumuitslag moet middels een partij- of batchnummer blijken dat de uitslag gerelateerd is aan de te exporteren partij. De uitslag mag maximaal vier weken oud zijn.

Voor een product vervaardigd in een andere EU-lidstaat kan deze verklaring worden afgegeven op basis van een veterinaire verklaring van gelijke strekking of door het overleggen van uitslagen van microbiologisch onderzoek op basis van partijbemonstering in het Nederlandse opslagbedrijf.

Voor een product vervaardigd in een derde land kan deze verklaring worden afgegeven door het overleggen van het EU-importcertificaat met daarin verklaringen van gelijke strekking plus bijbehorend GGB of door het overleggen van uitslagen van microbiologisch onderzoek op basis van partijbemonstering in het Nederlandse opslagbedrijf.

Verklaring II.5:

Have undergone all precautions to avoid contamination with pathogenic agents after treatment;
Deze verklaring kan worden afgegeven op basis van EU- en nationale regelgeving voor een product vervaardigd in een bedrijf met een registratie of erkenning op basis van Verordening (EG) nr. 1069/2009 of afkomstig van een bedrijf met een registratie op basis van Verordening (EG) nr. 183/2005.

Verklaring II.6:

Was packed in new packaging, which, if the pet food is not dispatched in ready-to-sell packages on which it is clearly indicated that the content is destined for feeding to pets only, bears labels indicating "NOT FOR HUMAN CONSUMPTION";

Deze verklaring kan worden afgegeven na controle. Bij invoeren van de aanvraag in e-CertNL moet het bedrijf aangeven/verklaren dat het product is verpakt in nieuwe verpakkingen. Voor bulkgoederen moet belanghebbende bovendien aantonen dat de verpakkingen zijn voorzien van een label met aanduiding: destined for feeding to pets only en "NOT FOR HUMAN CONSUMPTION".

Verklaring II.7:

The petfood described above⁽¹⁾

either⁽¹⁾ Is derived from other ruminants than bovine, ovine or caprine animals;

or⁽¹⁾ Is derived from bovine, ovine or caprine animals and does not contain and is not derived from:

either⁽¹⁾ Bovine, ovine or caprine materials other than those derived from animals born, continuously reared and slaughtered in a country or region which is classified as a "negligible" risk status by WOAH;

or⁽¹⁾ (a) Specified risk material as defined in point 1 of Annex V to Regulation (EC) No 999/2001 of the European Parliament and of the Council;

(b) Mechanically separated meat obtained from bones of bovine, ovine or caprine animals, except from those animals that were born, continuously reared and slaughtered in a country or region classified as posing a negligible BSE risk in accordance with WOAH, in which there has been no 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 WOAH.

Deze verklaring kan worden afgegeven na controle. De niet-relevante deelverklaringen moeten worden doorgehaald. Belanghebbende moet aantonen van welke grondstoffen het petfood is gemaakt (diersoort en herkomst). Deze verklaring is alleen van toepassing voor herkauwergondstoffen verkregen van het slachtproces en niet voor andere herkauwergondstoffen zoals zuivelgrondstoffen. Indien geen andere herkauwergondstoffen dan afkomstig van zuivel zijn verwerkt, moet de gehele verklaring worden doorgehaald.

Voor een product vervaardigd in Nederland of in een andere EU-lidstaat kan betreffende deelverklaring worden afgegeven op basis van een verklaring van gelijke strekking opgesteld door het productiebedrijf.

Voor een product vervaardigd in een derde land kan betreffende deelverklaring worden afgegeven door het overleggen van het EU-importcertificaat met daarin een verklaring van gelijke strekking plus bijbehorend GGB.

Voor een product vervaardigd in Nederland of in een andere EU-lidstaat geldt dat de items (a), (b) en (c) van de tweede deelverklaring kunnen worden afgegeven op basis van EU- en nationale regelgeving.

4.3 Dogchews

Certificaat: zie bijlage 3.

Algemeen:

- Raadpleeg vooraf de instructie Tijdelijke Maatregelen Derde Landen (TMDL-01) op mogelijke exportbeperkingen. Als in de TMDL-01 informatie staat die in strijd is met een landeninstructie dan is de informatie vermeld in de TMDL-01 leidend.
- Indien belanghebbende de gevraagde garanties niet kan aanleveren, zal de certificering niet doorgaan.

Invuleisen:

- I.8.: Dit veld hoeft alleen te worden ingevuld indien er sprake is van regionalisatie; zolang dit niet aan de orde is wordt er standaard de ISO-code van het land ingevuld.
- I.11.: In het certificaat kan slechts één producent worden opgenomen. Alleen producten van de betreffende producent kunnen op één certificaat gecertificeerd worden.
- I.19.: De HS-code voor kauwartinikelen is 2309. Alleen kauwartinikelen die onder deze code vallen mogen worden gecertificeerd. (In de voetnoot staat nog een HS-code, maar aangezien het certificaat producten van deze HS-code niet toestaat, is deze dan ook niet van toepassing).
- I.28.: In geval er meerdere soorten dierlijke grondstoffen in het product zijn verwerkt, dan moet de diersoort op grondstofniveau ingevuld worden, om te kunnen tonen op het certificaat.

Toelichting bij het certificaat:

Aanhef:

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 and in particular Article 10 of that Regulation, and Commission Regulation (EU) No 142/2011, and in particular Chapter II of Annex XIII and Chapter II of Annex XIV thereto, and certify that the dogchews described above:

Deze verklaring kan worden afgegeven indien de certificerende NVWA-dierenarts kennis heeft genomen van de relevante bepalingen van de genoemde EU-regelgeving (Verordening (EG) nr. 1069/2009 en Verordening (EG) nr. 142/2011).

Verklaring II.1:

have been prepared exclusively with the following animal by-products:

either⁽¹⁾ carcasses and parts of animal slaughtered or, in the case of game, bodies or parts of animal killed, and which are fit for human consumption, but are not intended for human consumption for commercial reasons;

and/or⁽¹⁾ carcasses 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:

i) carcasses or bodies and parts of animals which are rejected as unfit for human consumption, 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;

and/or⁽¹⁾ blood of animals which did not show any signs of disease communicable through blood to humans or animals, obtained from animals that have been slaughtered in a slaughterhouse after having been considered fit for slaughter for human consumption following an ante-mortem inspection;

and/or⁽¹⁾ animal by-products arising from the production of products intended for human consumption, including degreased bone, greaves and centrifuge or separator sludge from milk processing;

and/or⁽¹⁾ aquatic animals, and parts of such animals, except sea mammals, which did not show any signs of diseases communicable to humans or animals;

and/or⁽¹⁾ animal by-products from aquatic animals originating from plants or establishments manufacturing products for human consumption;
and/or⁽¹⁾ material from animals which have been treated with certain substances which are prohibited pursuant to Directive 96/22/EC, the import of the material being permitted in accordance with Article 35(a)(ii) of Regulation (EC) No 1069/2009;

Deze verklaring kan worden afgegeven op basis van een grondstoffenlijst. Belanghebbende moet de herkomst van de grondstoffen en de aard van het categorie 3-materiaal aantonen. De niet van toepassing zijnde opties kunnen worden doorgehaald.

De aanvrager dient zelf in de aanvraag in e-CertNL onder het tabblad "Documenten" bij "Verklaringsteksten" de keuze te maken welke tekst getoond wordt.

Voor een product vervaardigd in Nederland of in een andere EU-lidstaat kan deze verklaring worden afgegeven op basis van een grondstoffenlijst opgesteld door het productiebedrijf.

Voor een product vervaardigd in een derde land kan deze verklaring worden afgegeven door het overleggen van het EU-importcertificaat met daarin een verklaring van gelijke strekking plus bijbehorend GGB.

Verklaring II.2:

have been subjected:

either⁽¹⁾ in the case of dog chews made from hides and skins of ungulates or from fish, to a treatment sufficient to destroy pathogenic organisms (including salmonella); and the dog chews are dry;

and/or⁽¹⁾ in the case of dog chews made from animal by-products other than hides and skins of ungulates or from fish, to a heat treatment of at least 90 °C throughout their substance;

Deze verklaring kan worden afgegeven na controle. Belanghebbende moet aantoonbaar maken welke behandeling de te exporteren producten hebben ondergaan. Indien een behandeling niet van toepassing is, moet deze worden doorgehaald.

Voor de eerste optie geldt:

Voor dogchews van hoefdierhuiden of van vis die zijn vervaardigd in Nederland en andere EU-lidstaten kan dit op basis van een verklaring van gelijke strekking opgesteld door het productiebedrijf.

Voor dogchews vervaardigd in derde landen kan dit door het overleggen van het EU-importcertificaat met daarin een verklaring van gelijke strekking plus bijbehorend GGB.

Voor de tweede optie geldt:

Voor dogchews van andere grondstoffen die zijn vervaardigd in Nederland kan dit op basis van een bedrijfsverklaring aangaande het proces, in combinatie met periodieke verificatie.

Voor dogchews van andere grondstoffen die zijn vervaardigd in een andere EU-lidstaat kan dit op basis van een veterinaire verklaring van gelijke strekking. Dit kan zijn in de vorm van een certificaat bij iedere ontvangen zending, dan wel een 'long term declaration' (maximaal twaalf maanden oud).

Voor dogchews vervaardigd in derde landen kan dit door het overleggen van het EU-importcertificaat met daarin een verklaring van gelijke strekking plus bijbehorend GGB.

Verklaring II.3:

were examined by a random sampling of at least five samples from each processed batch taken during or after storage at the processing plant and complies with the following standards(2):

Salmonella : absence in 25 g: n = 5, c = 0, m = 0, M = 0

Enterobacteriaceae: n = 5, c = 2, m = 10, M=300 in 1 gram;

Deze verklaring kan worden afgegeven op basis van een laboratoriumuitslag voor de genoemde pathogenen, aan te leveren door de belanghebbende. Dit mag als volgt worden geïnterpreteerd: per certificaataanvraag moet één product per productiebedrijf zijn onderzocht (vijf deelmonsters, waarbij Salmonella door het laboratorium als mengmonster mag worden onderzocht). Uit de laboratoriumuitslag moet middels een partij- of batchnummer blijken dat de uitslag gerelateerd is aan de te exporteren partij. De uitslag mag maximaal vier weken oud zijn.

Voor een product vervaardigd in een andere EU-lidstaat kan deze verklaring worden afgegeven op basis van een veterinaire verklaring van gelijke strekking of door het overleggen van uitslagen van microbiologisch onderzoek op basis van partijbemonstering in het Nederlandse opslagbedrijf.

Voor een product vervaardigd in een derde land kan deze verklaring worden afgegeven door het overleggen van het EU-importcertificaat met daarin verklaringen van gelijke strekking plus bijbehorend GGB of door het overleggen van uitslagen van microbiologisch onderzoek op basis van partijbemonstering in het Nederlandse opslagbedrijf.

Verklaring II.4:

have undergone all precautions to avoid contamination with pathogenic agents after treatment;
Deze verklaring kan worden afgegeven op basis van EU- en nationale regelgeving voor een product afkomstig van een bedrijf met een registratie of erkenning op basis van Verordening (EG) nr. 1069/2009 of afkomstig van een bedrijf met een registratie op basis van Verordening (EG) nr. 183/2005.

Verklaring II.5:

were packed in new packaging;

Deze verklaring kan worden afgegeven na controle. Bij invoeren van de aanvraag in e-CertNL moet het bedrijf aangeven dat het product is verpakt in nieuwe verpakkingen

Verklaring II.6:

The dogchews described above⁽¹⁾

either⁽¹⁾ is derived from other ruminants than bovine, ovine or caprine animals;

or⁽¹⁾ is derived from bovine, ovine or caprine animals and does not contain and is not derived from:

- either⁽¹⁾ bovine, ovine or 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 WOAH;*
- or⁽¹⁾*
 - (a) specified risk material as defined in point 1 of Annex V to Regulation (EC) No 999/2001 of the European Parliament and of the Council;*
 - (b) mechanically separated meat obtained from bones of bovine, ovine or caprine animals, except from those animals that were born, continuously reared and slaughtered in a country or region classified as posing a negligible BSE risk in accordance with WOAH, in which there has been no 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 WOAH.*

Deze verklaring kan worden afgegeven na controle. De niet-relevante deelverklaringen moeten worden doorgedaald. Belanghebbende moet aantonen van welke grondstoffen de dogchews zijn gemaakt (diersoort en herkomst). Deze verklaring is alleen van toepassing voor herkauwergrondstoffen verkregen van het slachtproces en niet voor andere herkauwergrondstoffen zoals zuivelgrondstoffen. Indien geen andere herkauwergrondstoffen dan afkomstig van zuivel zijn verwerkt, moet de gehele verklaring worden doorgedaald.

Voor een product vervaardigd in Nederland of in een andere EU-lidstaat kan betreffende deelverklaring worden afgegeven op basis van een verklaring van gelijke strekking opgesteld door het productiebedrijf.

Voor een product vervaardigd in een derde land kan betreffende deelverklaring worden afgegeven door het overleggen van het EU-importcertificaat met daarin een verklaring van gelijke strekking plus bijbehorend GGB.

Voor een product vervaardigd in Nederland of in een andere EU-lidstaat geldt dat de items (a), (b) en (c) van de tweede deelverklaring kunnen worden afgegeven op basis van EU- en nationale regelgeving.

4.4 Smaakgevende ingewanden.

Voor export van smaakgevende ingewanden moet het certificaat voor processed petfood other than canned petfood worden aangevraagd. Zie paragraaf 4.2.

5 Bevoegdheden en verantwoordelijkheden

De NVWA-dierenarts is bevoegd en verantwoordelijk voor het afgeven van het certificaat.

Bijlage 1: certificaat canned petfood

TÜRKİYE CUMHURİYETİ'NE KONSERVE EV VE SÜS HAYVANI YEMİ
İHRACATI İÇİN VETERİNER SAĞLIK SERTİFİKASI /
VETERINARY HEALTH CERTIFICATE FOR EXPORTATION OF CANNED PETFOOD
TO THE REPUBLIC OF TURKEY

BÖLÜM I: GÖNDERİLEN SEVKİYATIN DETAYLARI / PART I: DETAILS OF DISPATCHED CONSIGNMENT						
I.1. Gönderen / Consignor Adı / Name Adresi / Address Posta Kodu / Postal code Telefonu / Tel.No.				I.2. Sertifika referans numarası / Certificate reference number I.3. Merkezi Yetkili Makam / Central Competent Authority I.4. Yerel Yetkili Makam / Local Competent Authority		I.2.a XXXX XX
I.5. Alıcı / Consignee Adı / Name Adresi / Address Posta kodu / Postal code Telefonu / Tel.No				I.6 XXXXXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXXXXX		
I.7. Menşe ülkesi// Country of origin The Netherlands	ISO Kodu / ISO code NL	I.8. Menşe bölgesi / Region of origin 	Kodu / Code 	I.9. Varış ülkesi / Country of destination TÜRKİYE	ISO Kodu / ISO code TR	I.10. XXXXX XXXXX XXXXX XXXXX XXXX X
I.11. Orjin yeri / Place of origin Adı / Name Onay numarası / Approval number Adresi / Address				I.12. XXXXXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXXXXX		
I.13. Yükleme yeri / Place of loading				I.14. Yola çıkış tarihi / Date of departure XXXXXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXXXXX		
I.15. Nakliye aracı / Means of transport Uçak / Aeroplane <input type="checkbox"/> Gemi / <input type="checkbox"/> Tren vagonu/Railway <input type="checkbox"/> Ship wagon Karayola taşıtı / <input type="checkbox"/> Diğer / <input type="checkbox"/> Road vehicle Other Tanımlama / Identification: Belge referansları / Documentary references:				I.16. Türkiye'ye giriş VSKN / Entry BIP in Turkey I.17. XXXXXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXXXXX		
I.18. Malın tanımı / Description of commodity				I.19. Malın kodu (HS kodu) / Commodity code (HS code)		I.20. Miktar / Quantity
I.21. Ürünün sıcaklığı / Temperature of product Oda sıcaklığı / Ambient <input type="checkbox"/> Soğutulmuş / Chilled <input type="checkbox"/> Dondurulmuş / Frozen				I.22. Paket sayısı / Number of packages		
I.23. Konteynerin tanımlaması/mühür numarası / Identification of container/seal number				I.24 Paketleme tipi /		

		Type of packaging			
1.25. Malların sertifikalandırma amacı / Commodities certified for:					
Pet hayvan yemi / <input checked="" type="checkbox"/> Teknik kullanım / <input type="checkbox"/> Petfood Technical use					
I.26. XXXXXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXXXXX XX	1.27. Türkiye'ye ithalat ya da kabul amaçlı / <input checked="" type="checkbox"/> For import into Turkey				
1.28. Malların tanımlaması / Identification of the commodities					
Parti numarası / Batch number	Türü (Bilimsel adı) / Species (Scientific name)	Malın tanımı / Nature of commodity	İşletmelerin Onay numarası: Üretim tesisi / Approval no of establishments: Manufacturing plant	Paket sayısı / Number of packages	Net ağırlık / Net weight

Ben, aşağıda imzası bulunan resmi veteriner hekim, Avrupa Parlamentosu ve Konseyi'nin 1069/2009/EC No'lu Yönetmeliği ve özellikle 8. ve 10. Maddesini, Komisyon 142/2011/EU No'lu Yönetmeliği ve özellikle Ek XIII, Bölüm II ve Ek XIV, Bölüm II' sini okuduğumu ve anladığımı beyan eder ve yukarıda tanımlanan ev ve süs hayvani yemlerine ilişkin olarak aşağıdakileri onaylarım: / 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 and in particular Article 8 and 10 thereof, and Commission Regulation (EU) No 142/2011, and in particular Chapter II of Annex XIII and Chapter II of Annex XIV thereof and certify that the petfood described above:

- II.1 1069/2009/EC No'lu Yönetmeliğin 24 üncü Maddesine uygun olarak yetkili otorite tarafından onaylanan ve denetlenen bir işletme veya tesiste hazırlanmış ve depolanmıştır / has been prepared and stored in an establishment or plant approved and supervised by the competent authority in accordance with Article 24 of Regulation (EC) No 1069/2009;
- II.2 Yalnızca aşağıdaki hayvansal yan ürünlerle hazırlanmıştır: / has been prepared exclusively with the following animal by-products:
 - ya / insan tüketimine uygun olan, ancak ticari sebeplerle insan tüketimine sunulması amaçlanmayarak kesilen hayvanların karkas ve parçaları veya öldürülen av hayvanlarının gövdesi veya parçaları / carcasses and parts of animal slaughtered or, in the case of game, bodies or parts of animal killed, and which are fit for human consumption, but are not intended for human consumption for commercial reasons;
 - ve/veya bir kesimhanede kesilen ve antemortem bir muayeneyi takiben insan tüketimi için kesilmeye uygun bulunan hayvanlardan orijin alan karkaslar ve aşağıdaki parçaları / carcasses 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:
 - (i) İnsan tüketimine uygun olmadığı için reddedilen, ancak insanlar veya hayvanlara geçebilen hastalık belirtisi göstermeyen hayvanların karkasları veya gövdeleri ile bunların parçaları / carcasses or bodies and parts of animals which are rejected as unfit for human consumption, but which did not show any signs of disease communicable to humans or animals;
 - (ii) kanatlı kafaları / heads of poultry;
 - (iii) kırıntı ve parçaları dahil post ve derileri, boynuzları ve falanks, karpal ve merakarpal kemikleri, tarsal ve metatarsal kemikleri dahil ayakları /

		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)	domuz kolları / pig bristles;
	(v)	tüyüler / feathers;
ve/veya / and/or ⁽¹⁾		853/2004/EC sayılı Yönetmeliğin madde 1(3)(d)'nde belirtilen çiftlikte insanlara ve hayvanlara geçebilen bulaşıcı hastalık belirtisi göstermeyen, kesilen kümes hayvanları ve tavşanımsılardan elde edilen hayvansal yan ürünler / animal by products from poultry and lagomorphs slaughtered on the farm as referred to in Article 1(3)(d) of Regulation (EC) No 853/2004, which did not show any signs of disease communicable to humans or animals;
ve/veya / and/or ⁽¹⁾		antemortem bir muayeneyi takiben insan tüketimi için kesilmeye uygun bulunduktan sonra bir kesimhanede kesilmiş olan hayvanlardan elde edilen kan yolu ile insan veya hayvanlara bulaşabilecek herhangi bir hastalık belirtisi göstermeyen hayvanların kanları / blood of animals which did not show any signs of disease communicable through blood to humans or animals, obtained from animals that have been slaughtered in a slaughterhouse after having been considered fit for slaughter for human consumption following an ante- mortem inspection;
ve/veya / and/or ⁽¹⁾		yağı alınmış kemikler, don yağı tortusu ve sütün işlenmesi sonucu ortaya çıkan santrifüj veya seperatör tortuları dahil insan tüketimi amaçlı ürünlerin üretiminden kaynaklanan hayvansal yan ürünler / animal by- products arising from the production of products intended for human consumption, including degreased bone, greaves and centrifuge or separator sludge from milk processing;
ve/veya / and/or ⁽¹⁾		ticari sebeplerle insan tüketimine sunulması amaçlanmayan veya üretim veya paketleme hataları bulunan veya halk ve hayvan sağlığı için risk taşımayan diğer kusurları olan hayvansal orijinli ürünler veya hayvansal orijinli ürün içeren gıda maddeleri / products of animal origin or foodstuffs containing products of animal origin, which are no longer intended for human consumption for commercial reasons or due to problems of manufacturing or packaging defects or other defects from which no risk to public or animal health arise;
ve/veya / and/or ⁽¹⁾		ticari sebeplerle yem olarak kullanılması amaçlanmayan veya üretim veya paketleme hataları bulunan veya halk veya hayvan sağlığı için risk taşımayan diğer kusurları olan ev ve süs hayvanı yemi ve hayvansal orjinli yem veya hayvansal yan ürün veya türev ürünlerini içeren yem maddeleri / petfood and feedingstuffs of animal origin, or feedingstuffs containing animal by-products or derived products, which are no longer intended for feeding for commercial reasons or due to problems of manufacturing or packaging defects or other defects from which no risk to public or animal health arises;
ve/veya / and/or ⁽¹⁾		o ürün yoluyla insan veya hayvanlara geçebilen herhangi bir hastalığın belirtisini göstermeyen canlı hayvanlardan elde edilen kan, plasenta, yün, tüy, kıl, boynuz, toynak kesikleri ve çiğ süt / blood, placenta, wool, feathers, hair, horns, hoof cuts and raw milk originating from live animals that did not show signs of any disease communicable through that product to humans or animals;
ve/veya / and/or ⁽¹⁾		insan veya hayvanlara geçebilen herhangi bir hastalığın belirtisini göstermeyen, deniz memelileri hariç su hayvanları ve bu hayvanların parçaları / aquatic animals, and parts of such animals, except sea mammals, which did not show any signs of diseases communicable to humans or animals;
ve/veya / and/or ⁽¹⁾		insan tüketimi için ürünler üreten işletme veya tesislerdeki su hayvanlarından elde edilen hayvansal yan ürünler / animal by-products from aquatic animals originating from plants or establishments manufacturing products for human consumption;
ve/veya / and/or ⁽¹⁾		o materyal yoluyla insan veya hayvanlara geçebilen herhangi bir hastalık belirtisi göstermeyen hayvanlardan elde edilen aşağıdaki ürünler : the following material originating from animals which did not show any signs of disease communicable through that material to humans or animals:
	i)	yumuşak doku veya etini içeren su kabuklarının kabukları / shells from shellfish with soft tissue or flesh;

- ii) kara hayvanlarından elde edilen aşağıdaki ürünler: /
the following originating from terrestrial animals;
 - kuluçka yan ürünleri /
hatchery by-products;
 - yumurta / eggs;
 - yumurta kabuğu içeren yumurta yan ürünleri / egg by-products, including egg shells;
 - iii) ticari sebeplerle öldürülen günlük civcivler / day-old chicks killed for commercial reasons;
- ve/veya
/ and/or⁽¹⁾ insan veya hayvanlar için spesifik patojen olanlar dışındaki su veya kara omurgasızları yan ürünleri / animal by-products from aquatic or terrestrial invertebrates other than species pathogenic to humans or animals;
- ve/veya
/ and/or⁽¹⁾ 1069/2009/EC sayılı Yönetmelik'in Madde 8(a)(iii), (iv) ve (v)'inde atıfta bulunulan Kategori 1 materyali ve Madde 9(a) ilâ (g)'sında atıfta bulunulan Kategori 2 materyali haricinde, Rodentia ve Lagomorpha zoojik düzenlerinden hayvanlar ve onları kısımları / animals and parts thereof of the zoological orders of Rodentia and Lagomorpha, except Category 1 material as referred to in Article 8(a)(iii), (iv) and (v) of Regulation (EC) No 1069/2009 and Category 2 material as referred to in Article 9(a) to (g) of that Regulation;
- ve/veya
/ and/or⁽¹⁾ 96/22/EC nolu direktife istinaden yasaklı olan ve 1069/2009 No.lu Yönetmeliği (EC) Bölüm 35 (a) (ii)' ye uygun olarak malzemelerin ithaline izin verilen belirli maddeler ile muameleye tutulan hayvanlardan elde edilen malzemeler / material from animals which have been treated with certain substances which are prohibited pursuant to Directive 96/22/EC, the import of the material being permitted in accordance with Article 35(a)(ii) of Regulation (EC) No 1069/2009;
- II.3 Hava sızdırmaz konteynirlarda minimum 3 F_c değeriyle ısıl işleme tabi tutulmuştur / has been subjected to heat treatment to a minimum F_c value of 3 in hermetically sealed containers;
- II.4 Madde II.3.'te öngörüldüğü şekilde tüm sevkiyatın yeterli ısıl işlem görmesini sağlamak için laboratuvar teşhis yöntemleriyle her bir işlenmiş partinin en az beş numune tesadüfi örneklemeye metoduyla analiz edilmiştir / was analysed by a random sampling of at least five samples from each processed batch by laboratory diagnostic methods to ensure adequate heat treatment of the whole consignment as foreseen under point II.3;
- II.5 İşlem sonrasında patojenik ajanlarla kontaminasyonu engelleyecek tüm önlemler alınmıştır / has undergone all precautions to avoid contamination with pathologic agents after treatment;
- II.6 Yukarıda açıklanan pet hayvan yemleri⁽¹⁾ / the petfood described above⁽¹⁾
- ya / either⁽¹⁾ Sığır, koyun cinsi hayvanlar dışındaki ruminantlardan elde edilmiştir / is derived from other ruminants than bovine, ovine or caprine animals)
 - veya / or⁽¹⁾ Sığır, koyun ve keçi cinsi hayvanlardan elde edilmiştir ve aşağıdakileri içermez ve bunlardan türetilmemiştir / is derived from bovine, ovine or caprine animals and does not contain and is not derived from:
 - ya / either⁽¹⁾ Dünya Hayvan Sağlığı Teşkilatının (WOAH) sınıflandırmasına göre "ihmal edilebilir" BSE risk statüsünde yer alan bir ülkede veya bölgede doğmuş, sürekli olarak burada yetiştirilmiş ve kesilmiş hayvanlardan elde edilenler dışındaki sığır, koyun veya keçi cinsi hayvanların ürünlerini / bovine, ovine or caprine materials other than those derived from animals born, continuously reared and slaughtered in a country or region which is classified as a "negligible" risk status by WOAH;
 - veya / or⁽¹⁾ (a) Avrupa Konseyi ve Parlamentosu'nun 999/2001 (EC) sayılı Yönetmeliği'nin Ek V' in 1. maddesinde tanımlanan spesifik risk materyali / specified risk material as defined in point 1 of Annex V to Regulation (EC) No 999/2001 of the European Parliament and of the Council;
 - (b) Yöresel olarak BSE vakası bulunmayan, Dünya Hayvan Sağlığı Teşkilatı (WOAH) standartlarına göre ihmal edilebilir BSE risk statüsünde)

sınıflandırılmış bir ülkede veya bölgede sürekli olarak yetiştirilen ve kesilen sığır, koyun, keçi cinsi hayvanlar dışındaki hayvanların kemiklerinden mekanik olarak ayrılmış olan etleri / mechanically separated meat obtained from bones of bovine, ovine or caprine animals, except from those animals that were born, continuously reared and slaughtered in a country or region classified as posing a negligible BSE risk in accordance with WOAH, in which there has been no indigenous BSE case;

- (c) Dünya Hayvan Sağlığı Teşkilatı (WOAH) standartlarına göre ihmal edilebilir BSE risk statüsündeki ülke veya bölgede doğmuş, sürekli yetiştirilmiş ve kesilmiş hayvanlar dışında kafatası boşluğuna sokulmuş uzun çubuk şeklinde bir alet vasıtasiyla merkezi sinir sistemi dokularının harap edilmesi yoluyla veya kafatası boşluğuna gaz enjekte edilmesi yoluyla sersemletme sonrasında öldürülen sığır, koyun veya keçi cinsi hayvanlardan elde edilen hayvansal yan ürün ve türev ürünleri / 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 WOAH.

Notlar / Notes

Bölüm I / Part I:

- Kutu referansı I.15: Sicil numarası (tren vagonları veya konteynır ya da kamyonlar), uçuş sayısı (uçak) veya isim (gemi); bilgiler boşaltma veya yeniden yükleme halinde verilmelidir / Box reference 1.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship); information is to be provided in the event of unloading and reloading
- Kutu referansı I.23: Dökme yük konteynırlar için, konteynır numarası ve mühür numarası (varsayı) verilmelidir / Box reference I.23: for bulk containers, the container number and the seal number (if applicable) should be given
- Kutu referansı I.25: teknik kullanım: Çiftlik hayvanlarının, kürk hayvanlarının dışındaki hayvanların beslenmesi ve pet hayvan yemlerinin üretimi ya da imalatı için kullanılması / Box reference I.25: technical use: any use other than for feeding of farmed animals, other than fur animals, and the production or manufacturing of pet food..

Bölüm II / Part II:

- (1) Uygun şekilde silin /
-) Delete as appropriate
- İmza ve mühür renginin, baskından farklı bir renkte olması gerekmektedir / The signature and the stamp must be in a different colour to that of the printing
- Türkiye'de ki sevkiyattan sorumlu kişinin dikkatine: Bu sertifika yalnızca veteriner kullanım amaçlıdır ve ilgili sınır kontrol noktasına varana kadar sevkiyata eşlik etmelidir / Note for the person responsible for the consignment in TR: this certificate is only for veterinary purposes and has to accompany the consignment until it reaches the border inspection post.

Bijlage 2: certificaat other than canned petfood

**TÜRKİYE CUMHURİYETİ'NE KONSERVE PET HAYVANI YEMİ DIŞINDAKİ İŞLENMİŞ PET HAYVANI YEMİ İHRACATI İÇİN VETERİNER SAĞLIK SERTİFİKASI /
VETERINARY HEALTH CERTIFICATE FOR EXPORTATION OF PROCESSED PETFOOD OTHER THAN
CANNED PETFOOD TO THE REPUBLIC OF TURKEY**

BÖLÜM I: GÖNDERİLEN SEVKIYATIN DETAYLARI / PART I: DETAILS OF DISPATCHED CONSIGNMENT						
I.1. Gönderen / Consignor Adı / Name Adresi / Address Posta Kodu / Postal code Telefonu / Tel.No.				I.2. Sertifika referans numarası / Certificate reference number XXXXXXXX	I.2.a I.3 Merkezi Yetkili Makam / Central Competent Authority XXXXXXXXXXXXXXXXXXXXXXXXX	
				I.4. Yerel Yetkili Makam / Local Competent Authority XXXXXXXXXXXXXXXXXXXXXXXXX		
I.5. Alıcı / Consignee Adı / Name Adresi / Address Posta kodu / Postal code Telefonu / Tel.No				I.6 XXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXX		
I.7. Menşe ülkesi/ Country of origin The Netherlands	ISO Kodu / ISO code NL	I.8. Menşe bölgesi / Region of origin XXXXXXXXX	Kodu / Code TÜRKİYE	I.9. Varış ülkesi / Country of destination TÜRKİYE	ISO Kodu / ISO code TR	I.10. XXXX XXXX XXXX XXXX XXXX XXXX XXXX XXXX
I.11. Orjin yeri / Place of origin Adı / Name Onay numarası / Approval number Adresi / Address				I.12. XXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXX		
I.13. Yükleme yeri / Place of loading Uçak / Aeroplane <input type="checkbox"/> Gemi / Ship <input type="checkbox"/> Tren vagonu/Railway wagon <input type="checkbox"/> Karayola taşıtı / <input type="checkbox"/> Diğer / Other <input type="checkbox"/>				I.14. Yola çıkış tarihi / Date of departure I.16. Türkiye'ye giriş VSKN / Entry BIP in Turkey XXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXX		
Tanımlama / Identification: Belge referansları / Documentary references:				I.17. XXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXX		
I.18. Malın tanımı / Description of commodity				I.19. Malın kodu (HS kodu) / Commodity code (HS code) I.20. Miktar / Quantitiy		
I.21. Ürünün sıcaklığı / Temperature of product Oda sıcaklığı / <input type="checkbox"/> Soğutulmuş / Chilled <input type="checkbox"/> Dondurulmuş / Frozen <input type="checkbox"/>				I.22. Paket sayısı / Number of packages		

I.23. Konteynerin tanimlamasi/mühür numarası / Identification of container/seal number	I.24 Paketleme tipi / Type of packages				
1.25. Malların sertifikalandırma amacı / Commodities certified for: Pet hayvan yemi / <input checked="" type="checkbox"/> Teknik kullanım / <input type="checkbox"/> Petfood Technical use					
I.26. XXXXXXXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXXXXXXX	1.27. Türkiye'ye ithalat ya da kabul amaçlı / <input checked="" type="checkbox"/> For import into Turkey				
1.28. Malların tanımlaması / Identification of the commodities					
Parti numarası / Batch number	Türü (Bilimsel adı) / Species (Scientific name)	Malın tanımı / Nature of commodity	İşletmelerin Onay numarası: Üretim tesisi / Approval no of establishments: Manufacturing plant	Paket sayısı / Number of packages	Net ağırlık / Net weight

Ben, aşağıda imzası bulunan resmi veteriner hekim, Avrupa Parlamentosu ve Konseyi'nin 1069/2009/EC No'lu Yönetmeliği ve özellikle 8. ve 10. Maddesini, Komisyon 142/2011/EU No'lu Yönetmeliği ve özellikle Ek XIII, Bölüm II ve Ek XIV, Bölüm II' sini okuduğumu ve anladığımı beyan eder ve yukarıda tanımlanan ev ve süs hayvanı yemlerine ilişkin olarak aşağıdakileri onaylarım: / 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 and in particular Articles 8 and 10 thereof, and Commission Regulation (EU) No 142/2011, and in particular Chapter II of Annex XIII and Chapter II of Annex XIV thereto and certify that the petfood described above:

- II.1. 1069/2009 No'lu Yönetmeliğin (EC) 24 üncü Maddesine uygun olarak yetkili otorite tarafından onaylanan ve denetlenen bir tesiste hazırlanmış ve depolanmıştır / 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;
- II.2. Yalnızca aşağıdaki hayvansal yan ürünlerle hazırlanmıştır: / Has been prepared exclusively with the following animal by-products:
 ya / Insan tüketimine uygun olan, ancak ticari sebeplerle insan tüketimine sunulması amaçlanmayarak kesilen hayvanların karkas ve parçaları veya öldürülen av hayvanlarının gövdesi veya parçaları / Carcasses and parts of animal slaughtered or, in the case of game, bodies or parts of animal killed, and which are fit for human consumption, but are not intended for human consumption for commercial reasons;
 ve/veya⁽¹⁾ Bir kesimhanede kesilen ve antemortem bir muayeneyi takiben insan tüketimi için kesilmeye uygun bulunan hayvanlardan orjin alan karkaslar ve aşağıdaki parçaları veya insan tüketimi için öldürülen av hayvanlarının gövde ve aşağıdaki parçaları: / Carcasses 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:
 i) insan tüketimine uygun olmadığı için reddedilen, ancak insanlar veya hayvanlara geçebilen hastalık belirtisi göstermeyen hayvanların karkasları veya gövdeleri ile bunların parçaları / carcasses or bodies and parts of animals which are rejected as unfit for human consumption, but which did not show any signs of disease communicable to humans or animals;
 ii) kanatlı kafaları / heads of poultry ;

	iii)	kırıntı ve parçaları dahil post ve derileri, boynuzları ve falanks, karpal ve merakarpal kemikleri, tarsal ve metatarsal kemikleri dahil ayakları / 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)	domuz kilları /pig bristles;
	v)	tüyler / feathers;
ve/veya ¹⁾ / and/or	853/2004/EC sayılı Yönetmeliğin madde 1(3)(d)'nde belirtilen çiftlikte insanlara ve hayvanlara geçebilen bulaşıcı hastalık belirtisi göstermeyen, kesilen kümes hayvanları ve tavşanımsılardan elde edilen hayvansal yan ürünler / Animal by products from poultry and lagomorphs slaughtered on the farm as referred to in Article 1(3)(d) of Regulation (EC) No 853/2004, which did not show any signs of disease communicable to humans or animals;	
ve/veya ¹⁾ / and/or	Antemortem bir muayeneyi takiben insan tüketimi için kesilmeye uygun bulunduktan sonra bir kesimhanede kesilmiş olan hayvanlardan elde edilen kan yolu ile insan veya hayvanlara bulaşabilecek herhangi bir hastalık belirtisi göstermeyen hayvanların kanları / Blood of animals which did not show any signs of disease communicable through blood to humans or animals, obtained from animals that have been slaughtered in a slaughterhouse after having been considered fit for slaughter for human consumption following an ante- mortem inspection;	
ve/veya ¹⁾ / and/or	Yağı alılmış kemikler, don yağı tortusu ve sütün işlenmesi sonucu ortaya çıkan santrifüj veya seperatör tortuları dahil insan tüketimi amaçlı ürünlerin üretiminde kaynaklanan hayvansal yan ürünler / Animal by- products arising from the production of products intended for human consumption, including degreased bone, greaves and centrifuge or separator sludge from milk processing;	
ve/veya ¹⁾ / and/or	Ticari sebeplerle yem olarak kullanılması amaçlanmayan veya üretim veya paketleme hataları bulunan veya halk veya hayvan sağlığı için risk taşımayan diğer kusurları olan ev ve süs hayvanı yemi ve hayvansal orjinli yem veya hayvansal yan ürün veya türev ürünlerini içeren yem maddeleri / Petfood and feedingstuffs of animal origin, or feedingstuffs containing animal by-products or derived products, which are no longer intended for feeding for commercial reasons or due to problems of manufacturing or packaging defects or other defects from which no risk to public or animal health arises;	
ve/veya ¹⁾ / and/or	O ürün yoluyla insan veya hayvanlara geçebilen herhangi bir hastalığın belirtisini göstermeyen canlı hayvanlardan elde edilen kan, plasenta, yün, tüy, kıl, boynuz, toynak kesikleri ve çiğ süt / Blood, placenta, wool, feathers, hair, horns, hoof cuts and raw milk originating from live animals that did not show signs of any disease communicable through that product to humans or animals;	
ve/veya ¹⁾ / and/or	İnsan veya hayvanlara geçebilen herhangi bir hastalığın belirtisini göstermeyen deniz memelileri hariç su hayvanları ve bu hayvanların parçaları / Aquatic animals, and parts of such animals, except sea mammals, which did not show any signs of diseases communicable to humans or animals;	
ve/veya ¹⁾ / and/or	İnsan tüketimi için ürünler üreten işletme veya tesislerdeki su hayvanlarından elde edilen hayvansal yan ürünler / Animal by-products from aquatic animals originating from plants or establishments manufacturing products for human consumption;	
ve/veya ¹⁾ / and/or	O materyal yoluyla insan veya hayvanlara geçebilen herhangi bir hastalık belirtisi göstermeyen hayvanlardan elde edilen aşağıdaki ürünler : The following material originating from animals which did not show any signs of disease communicable through that material to humans or animals:	
	i)	yumuşak doku veya etini içeren su kabuklarının kabukları / shells from shellfish with soft tissue or flesh;
	ii)	kara hayvanlarından elde edilen aşağıdaki ürünler: / the following originating from terrestrial animals:
		- kuluçka yan ürünleri / hatchery by-products,
		- yumurta /eggs;

			- yumurta kabuğu içeren yumurta yan ürünleri / egg by-products, including egg shells;
		iii)	ticari sebeplerle öldürülen günlük civcivler / day-old chicks killed for commercial reasons;
	ve/veya ⁽¹⁾ / and/or	Insan veya hayvanlar için spesifik patojen olanlar dışındaki su veya kara omurgasızları yan ürünleri / Animal by-products from aquatic or terrestrial invertebrates other than species pathogenic to humans or animals;	
	ve/veya ⁽¹⁾ / and/or	1069/2009/EC sayılı Yönetmelik'in Madde 8(a)(iii), (iv) ve (v)'inde atıfta bulunulan Kategori 1 materyali ve Madde 9(a) ilâ (g)'sında atıfta bulunulan Kategori 2 materyali haricinde, Rodentia ve Lagomorpha zoologik düzenlerinden hayvanlar ve onları kısımları / Animals and parts thereof of the zoological orders of Rodentia and Lagomorpha, except Category 1 material as referred to in Article 8(a)(iii), (iv) and (v) of Regulation (EC) No 1069/2009 and Category 2 material as referred to in Article 9(a) to (g) of that Regulation;	
	ve/veya ⁽¹⁾ / and/or	96/22/EC nolu direktifte istinaden yasaklı olan ve 1069/2009 No.lu Yönetmeliği (EC) Bölüm 35 (a) (ii)' ye uygun olarak malzemelerin ithaline izin verilen belirli maddeler ile muameleye tutulan hayvanlardan elde edilen malzemeler / Material from animals which have been treated with certain substances which are prohibited pursuant to Directive 96/22/EC, the import of the material being permitted in accordance with Article 35(a)(ii) of Regulation (EC) No 1069/2009;	
II.3.	ya / either ⁽¹⁾ ya da / or ⁽¹⁾	Madde genelinde en az 90°C'lik bir ısıl işleme tabi tutulmuştur / Was subjected to a heat treatment of at least 90°C throughout its substance; Yalnızca aşağıdaki koşulları sağlayan ürünler kullanılarak, hayvansal kökenli içerikler göz önünde bulundurularak üretilmiştir: / Was produced as regards ingredients of animal origin using exclusively products which had been: a. Madde genelinde en az 90°C'lik bir ısıl işleme tabi tutulmuş hayvansal yan ürünler veya et veya et ürünlerinden elde edilen türev ürünler olması durumunda / In the case of animal by- products or derived products from meat or meat products subjected to a heat treatment of at least 90°C throughout its substance; b. Süt ve süt kaynaklı ürünler olması durumunda / In the case of milk and milk based products, i) Şap hastalığından arı bir ülkeden/ bölgeden gelmiş ve içinde kullanılan çiğ süt, en az 72°C'de 15 saniyelik bir pastörizasyon işleminden elde edilene eşdeğer ısı etkisi yapacak bir işlemenin geçmiş ve phosphatase testinden negatif sonuç alınmıştır / Coming from country/region where are free from foot-and-mouth disease; and raw milk used in the product was treated with a heating effect at least equivalent to that achieved by a pasteurisation process of at least 72°C for 15 seconds to produce a negative phosphatase test;	
	ya / or ⁽¹⁾ ii)	Son 12 aydır şap hastalığı görülmeyen ülkeden/bölgeden gelmektedir ve içinde kullanılan çiğ sütün pH değeri 6'nın altına düşürülmüş ve ilk olarak yeterli miktarda bir pastörizasyon işlemi uygulanmış ve phosphatase testinden negatif sonuç alınmıştır / Coming from a country/region where has not been occurred any foot-and-mouth disease for last 12 months and pH of raw milk used in the product was reduced to less than 6 and firstly submitted to a pasteurisation treatment sufficient to produce a negative phosphatase test;	
	ya / or ⁽¹⁾ iii)	Son 12 aydır şap hastalığı görülmeyen ülkeden/bölgeden gelmektedir ve içinde kullanılan çiğ süte bir sterilizasyon işlemi ya da her bir işlem için yeterli miktarda çift ısı işlemi uygulanmış ve phosphatase testinden negatif sonuç alınmıştır / Coming from a country/region where has not been occurred any foot-and-mouth disease for last 12 months and raw milk used in the product was submitted to a sterilisation process or a double	

- heat treatment where each treatment was sufficient to produce a negative phosphatase test;
- ya / or⁽¹⁾ iv) Son 12 ay içinde şap hastalığı vakası görülmüş ya da son 12 aydır şap hastalığına karşı aşılama uygulanan ülkeden/bölgeden aşağıdaki işlemleri takiben gelmektedir: /
 Coming from a country where has been occurred foot-and-mouth disease for last 12 months or where vaccination against foot-and-mouth disease has been carried out in the last 12 months submitted to:
- ya da / En az 3 F_c değerine eşdeğer veya daha büyük sağlanan either bir sterilizasyon işlemi /
 A sterilisation process whereby an F_c value equal or greater than 3 is achieved;
- ya / or En az 72°C'de en az 15 saniyelik ve yeterli miktarda bir pastörizasyon işleminden elde edilene eşdeğer ısı etkisi yapacak bir başlangıç ısıl işlemi ve phosphatase testinden negatif sonuç alınmasını takiben: /
 An initial heat treatment with a heating effect at least equal to that achieved by a pasteurisation process of at least 72°C for at least 15 seconds and sufficient to produce a negative reaction to a phosphatase test, followed by:
- ya / Kurutulmuş süt ya da kurutulmuş süt kaynaklı ürünler either olması durumunda kurutma işleminden önce başlangıç ısıl işleminden elde edilene en az eşdeğer bir ısı etkisi yapacak ve phosphatase testine negatif reaksiyon verecek yeterlikte uygulanan ikinci bir ısıl işlem /
 A second heat treatment with a heating effect at least equal to that achieved by the initial heat treatment, and which would be sufficient to produce a negative reaction to a phosphatase test, followed, in the case of dried milk, or dried milk-based products by a drying process;
- ya da / En az 1 saat süresince pH değeri 6'nın altında olacak or şekilde bir asidifikasyon işlemi uygulanmıştır /
 An acidification process such that the pH has been maintained at less than 6 for at least one hour;
- c. Jelatin durumunda, işlenmemiş Kategori 3 materyalin asit veya alkaliyle bir işleme tabi tutulmasını sağlayan bir proses kullanılarak üretilmiş, bunu takiben pH ve daha sonrasının ayarlanmasıyla bir veya daha fazla durulama yapılmış, tekrarının gerektiği durumda, ısıtmaya ekstraksiyon yapılarak, bunu takiben filtrasyon ve sterilizasyon yoluyla arıtma yapılan /
 In the case of gelatine, produced using a process that ensures that unprocessed Category 3 material is subjected to a treatment with acid or alkali, followed by one or more rinses with subsequent adjustment of the pH and subsequent , if necessary repeated, extraction by heat, followed by purification by means of filtration and sterilisation;
- d. Kategori 3 materyalin kontaminasyonunu en aza indirmek için uygun önlemler içeren bir üretim prosesi kullanılarak üretilmiş hidrolize edilmiş protein durumunda, yalnızca hidrolize edilmiş protein üretimine ayrılmış bir işleme tesisisinde, ruminant post ve derilerinden tamamen veya kısmen üretilmiş hidrolize edilmiş protein olması durumunda, sadece 10.000 Dalton altında bir moleküller ağırlığa sahip malzeme kullanılarak ve Kategori 3 materyalin salamura etme, kirekleme ve yoğun yıkamıyla hazırlanmasını içeren bir proses kullanılarak, sonrasında aşağıdaki işlemler uygulanan: /
 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) Malzemenin 80°C'nin üzerinde bir sıcaklıkta üç saatten uzun süreyle 11'in üzerinde bir pH'a maruz bırakılması ve sonrasında 3,6 bar'ın üzerinde 30 dakika boyunca 140°C'nin üzerinde ıslı işleme tabi tutulması /
Exposure of the material to a pH 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;

veya/
or

ii) Malzemenin 1 ila 2'lik bir pH'a tabi tutulması, arkasından 11'in üzerinde bir pH'a tabi tutulması, arkasından 3 bar'da 30 dakika boyunca 140°C'lik bir ıslı işleme tabi tutulması /
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;

e. Yumurta ürünleri olması durumunda, 142/2011 Nolu Yönetmeliğin (EC) Ek IV Bölüm III'de dechinildiği gibi, 1'den 5'e kadar olan veya 7 nolu işleme yöntemlerinden herhangi birine tabi tutulan; veya Avrupa Parlamentosu ve Konseyi 853/2004 Nolu Yönetmeliğinin (EC) Ek III'nün X Kısmına ait Bölüm II'ye uygun olarak işlemden geçirilen / In the case of egg products submitted to any of the processing methods 1 to 5 or 7 as referred to in Chapter III of Annex IV to Regulation (EU) No 142/2011 or treated in accordance with Chapter II of Section X of Annex III to Regulation (EC) No 853/2004 of the European Parliament and of the Council;

f. Kollajen olması durumunda, AB Mevzuatınca izin verilenin dışında koruyucu kullanımını yasaklanmış, işlenmemiş kategori 3 materyalin yıkama, bir veya daha fazla kez durulanmasını takiben asit veya alkali kullanarak pH ayarlama, filtrasyon ve ekstrüzyon işlemlerinden geçirilen /

In the case of collagen submitted to a process ensuring that unprocessed Category 3 material is subjected to a treatment involving washing, pH adjustment using acid or alkali followed by one or more rinses, filtration and extrusion, the use of preservatives other than those permitted by Union legislation being prohibited;

g. Kan ürünleri olması durumunda, 142/2011 Nolu Yönetmeliğin (EC) Ek IV Bölüm III'de dechinildiği gibi, 1'den 5'e kadar olan veya 7 nolu işleme yöntemlerinden herhangi biri kullanılarak üretilen / In the case of blood products produced using any of the processing methods 1 to 5 or 7, as referred to in Chapter III of Annex IV to Regulation (EU) No 142/2011;

h. Memeli işlenmiş hayvan proteini durumunda, 1'den 5'e kadar olan veya 7 nolu işleme yöntemlerinden herhangi birine tabi tutulan ve, domuz kanı olması durumunda, 1'den 5'e kadar olan veya 7 nolu işleme yöntemlerinden herhangi birine tabi tutulan ancak 7 nolu yöntemin kullanılması durumunda madde genelinde en az 80°C sıcaklıkta bir ısı işlemi uygulanan /

In the case of mammalian processed animal protein submitted to any of the processing methods 1 to 5 or 7 and in the case of porcine blood, submitted to any of the processing methods 1 to 5 or 7 provided that in the case of method 7 a heat treatment throughout its substance at a minimum temperature of 80°C has been applied;

i. Balık unu hariç memeli olmayan işlenmiş protein olması durumunda, 142/2011 Nolu Yönetmeliğin (EC) Ek IV Bölüm III'de dechinildiği gibi, 1'den 5'e kadar olan veya 7 no.lu işleme yöntemlerinden herhangi birine tabi tutulan /

- j. In the case of non-mammalian processed protein with the exclusion of fishmeal submitted to any of the processing methods 1 to 5 or 7 as referred to in Chapter III of Annex IV to Regulation (EU) No 142/2011; Balık unu olması durumunda 142/2011 sayılı yönetmeliğin bölüm III Ek IV'te belirtildiği üzere 1'den 7'ye kadar işleme metoduna tabi tutulan veya türev ürünler için 142/2011 sayılı yönetmeliğin bölüm I Ek X'da belirtilen mikrobiyolojik standartları sağlaması için yöntem ve parametrelere tabi tutulan; /
In the case of fishmeal submitted to any of the processing methods or to a method and parameters which ensure that the products complies with the microbiological standards for derived products set in Chapter I of Annex X to Regulation (EU) No 142/2011;
- k. Balık yağları da dahil olmak üzere, rendering yağı olması durumunda, 142/2011 No.lu Yönetmeliğin (EC) Ek IV Bölüm III'de debynildiği gibi, 1'den 5'e kadar olan veya 7 nolu (ve balık yağı durumunda 6 nolu yöntem) işleme yöntemlerine tabi tutulan veya 853/2004 Nolu Yönetmeliğin (EC) Ek III'nün XII Kısımına ait Bölüm II'ye uygun olarak üretilen; ruminantlardan elde edilen rendering yağlarda, kalan toplam çözünmez katkıların maksimum seviyesi ağırlık olarak % 0,15'i aşmayacak şekilde arıtlan /
In the case of rendered fat including fish oils, submitted to any of the processing methods 1 to 5 or 7 (and method 6 in the case of fish oil) as referred to in Chapter III of Annex IV to Regulation (EU) No 142/2011 or produced in accordance with Chapter II of Section XII of Annex III to Regulation (EC) No 853/2004; rendered fats from ruminant animals must be purified in such a way that the maximum level of remaining total insoluble impurities does not exceed 0,15% in weight;
- l. Dikalsiyum fosfat olması durumunda üretimde aşağıda belirtilen işlemler uygulanır: /
In the case of dicalcium phosphate produced by a process that:
- i) Tüm kategori 3 kemik-materyalin ince şekilde ezilmesini ve sıcak suyla yağıdan arındırılmasını sağlayan ve en az iki günlük bir süre boyunca sulandırılmış hidroklorik asitle (en az %4 yoğunlukta ve 1,5'tan az bir pH ile) işleme tabi tutulmasını sağlayan /
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;
 - ve/ and ii) Madde (i) kapsamındaki işlemi takiben, elde edilen fosforik likörün kireçle muamele edilmesiyle 4 ila 7 pH'de dikalsiyum fosfat çökteltisi elde eden /
Following the procedure under (i), applied a treatment of the obtained phosphoric liquor with lime, resulting in a precipitate of dicalcium phosphate at pH 4 to 7;
 - iii) Son olarak, 65°C - 325°C'lik bir giriş sıcaklığı ve 30°C - 65°C 'lik bir bitiş sıcaklığında dikalsiyum fosfat çökteltisini havayla kurutan /
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. Trikalsiyum fosfat olması durumunda, aşağıdakileri sağlayan bir işlemle üretilen: /
In the case of tricalcium phosphate produced by a process that ensures:
- i) Tüm kategori 3 kemik-materyalin ince şekilde ezilmesini ve sıcak suyla ters akış uygulanarak yağıdan arındırılmasını sağlayan (kemik kıritülleri 14 mm'den az) /
That all Category 3 bone-material is finely crushed and degreased in counter-flow with hot water (bone chips less than 14 mm);
 - ii) 4 bar'da 30 dakika boyunca 145°C'de buharla sürekli pişirilen /

- Continuous cooking with steam at 145°C during 30 minutes at 4 bar;
- ve/ and n. iii) Protein besiyerinin santrifügasyon ile hidroksiapatitten (trikalsiyum fosfatat) ayrılmasını sağlayan / Separation of the protein broth from the hydroxyapatite (tricalcium phosphate) by centrifugation;
- ve/ and n. iv) 200°C'de hava akışkanlı bir yatak içinde kurutulduktan sonra trikalsiyum fosfatın granülasyonunu sağlayan / Granulation of the tricalcium phosphate after drying in a fluid bed with air at 200°C; İç organ Tatlandırıcı olması durumunda; ürünün madde II:4 de belirtilen mikrobiyolojik standartlara uygunluğunu sağlayan muamele yöntem ve parametrelere göre üretilen / In the case of flavouring innards, produced according to treatment method and parameters, which ensure that the product complies with the microbiological standards referred to under point II.4;
- ya / or⁽¹⁾ Yetkili otorite tarafından izin verilen kurutma veya fermantasyon gibi bir muameleye tabi tutulan / Was subject to a treatment such as drying or fermentation, which has been authorised by the competent authority;
- ya / or⁽¹⁾ İnsan veya hayvanlar için patojen türlerin dışında su ve kara omurgasız hayvanı olması durumunda, yetkili otorite tarafından izin verilen ve pet yeminin insan ve hayvan sağlığı için kabul edilemez hiçbir risk taşımadığını garanti eden bir muameleye tabi tutulan / In the case of aquatic and terrestrial invertebrates other species pathogenic to humans or animals, be subject to a treatment which has been authorised by the competent authority and ensures that the pet food poses no unacceptable risks to public and animal health;
- II.4. İşleme tesisinde depolama sırasında veya sonrasında her bir işlenmiş partiden alınan en az beş numune rastgele örnekleme yoluyla alınarak analiz edilmiş ve aşağıdaki standartlara⁽²⁾ uygun bulunmuştur: / Were analysed by a random sampling of at least five samples from each processed batch taken during or after storage at the processing plant and complies with the following standards⁽²⁾:
- Salmonella: 25 g lik örnekte hiç yok: n=5, c=0, M=0 / absence in 25 g: n = 5, c = 0, M = 0
 - Enterobacteriaceae: 1 gr lik örnekte en çok n=5, c=2, m=10, M=300 / n = 5, c = 2, m = 10, M=300 in 1 gram;
- II.5. İşlem sonrasında patojenik ajanlarla kontaminasyonu engelleyecek tüm önlemler alınmıştır / Have undergone all precautions to avoid contamination with pathogenic agents after treatment;
- II.6. Eğer ev ve süs hayvanı yemi, üzerinde yalnızca hayvanları besleme amaçlı olduğu hususunun açıkça belirtildiği satışa hazır paketlerde gönderilmiyorsa, "İNSAN TÜKETİMİ İÇİN DEĞİLDİR" ibaresinin yazılı olduğu etiketler taşıyan yeni paketlerde ambalajlanmıştır / Was packed in new packaging, which, if the pet food is not dispatched in ready-to-sell packages on which it is clearly indicated that the content is destined for feeding to pets only, bears labels indicating "NOT FOR HUMAN CONSUMPTION";
- II.7. Yukarıda açıklanan pet hayvan yemleri⁽¹⁾ / The petfood described above⁽¹⁾
- ya / either⁽¹⁾ Sığır, koyun veya keçi cinsi hayvanlar dışındaki ruminantlardan elde edilmiştir / Is derived from other ruminants than bovine, ovine or caprine animals;
- veya / or⁽¹⁾ Sığır, koyun veya keçi cinsi hayvanların ürünlerini içermemektedir ve bunlardan elde edilmemiştir: / Is derived from bovine, ovine or caprine animals and does not contain and is not derived from:
- ya / either⁽¹⁾ Dünya Hayvan Sağlığı Teşkilatının (WOAH) sınıflandırmasına göre "ihmal edilebilir" BSE risk statüsünde yer alan bir ülkede veya bölgede doğmuş, sürekli olarak burada yetiştirilmiş ve kesilmiş hayvanlardan elde edilenler dışındaki sığır, koyun veya keçi cinsi hayvanların ürünlerini / Bovine, ovine or caprine materials other than those derived from animals born, continuously reared and slaughtered in a country or region which is classified as a "negligible" risk status by WOAH;

- veya /
or⁽¹⁾
- (a) Avrupa Konseyi ve Parlamentosu'nun 999/2001 (EC) sayılı Yönetmeliği'nin Ek V' in 1. maddesinde tanımlanan spesifik risk materyali / Specified risk material as defined in point 1 of Annex V to Regulation (EC) No 999/2001 of the European Parliament and of the Council;
 - (b) Yöresel olarak BSE vakası bulunmayan, Dünya Hayvan Sağlığı Teşkilatı (WOAH) standartlarına göre ihmal edilebilir BSE risk statüsünde sınıflandırılmış bir ülkede veya bölgede sürekli olarak yetiştirilen ve kesilen sığır, koyun, keçi cinsi hayvanlar dışındaki hayvanların kemiklerinden mekanik olarak ayrılmış olan etleri / Mechanically separated meat obtained from bones of bovine, ovine or caprine animals, except from those animals that were born, continuously reared and slaughtered in a country or region classified as posing a negligible BSE risk in accordance with WOAH, in which there has been no indigenous BSE case;
 - (c) Dünya Hayvan Sağlığı Teşkilatı (WOAH) standartlarına göre ihmal edilebilir BSE risk statüsündeki ülke veya bölgede doğmuş, sürekli yetişтирilmiş ve kesilmiş hayvanlar dışında kafatası boşluğuna sokulmuş uzun çubuk şeklinde bir alet vasıtasyyla merkezi sinir sistemi dokularının harap edilmesi yoluyla veya kafatası boşluğuna gaz enjekte edilmesi yoluyla sersemletme sonrasında öldürülün sığır, koyun veya keçi cinsi hayvanlardan elde edilen hayvansal yan ürün ve türev ürünleri / 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 WOAH.

Notlar / Notes

Bölüm I / Part I:

- Kutu referansı I.15: Sicil numarası (tren vagonları veya konteynır ya da kamyonlar), uçuş sayısı (uçak) veya isim (gemi); bilgiler boşaltma veya yeniden yükleme halinde verilmelidir / Box reference 1.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship); information is to be provided in the event of unloading and reloading;
- Kutu referansı I.19: Uygun olan GTIP kodunu seçiniz. 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, 3503 veya 3504 / Box reference 1.19: 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, 3503 or 3504;
- Kutu referansı I.23: Dökme yük konteynırlar için, konteynır numarası ve mühür numarası (varsayı) verilmelidir./ Box reference I.23: for bulk containers, the container number and the seal number (if applicable) should be given;
- Kutu referansı I.25: teknik kullanım: Çiftlik hayvanlarının, kürk hayvanlarının dışındaki hayvanların beslenmesi ve pet hayvan yemlerinin üretimi ya da imalatı için kullanılması / Box reference I.25: technical use: any use other than for feeding of farmed animals, other than fur animals, and the production or manufacturing of pet food;
- Kutu referansı I.28: Türler: Uygun olanı seçiniz: kanatlı, Memeli, Ruminant, Pesca, Yumuşakça, Kabaklılar, Omurgasızlar / Box reference I.28: Aves, Mammalia, Ruminantia, Pesca, Mollusca, Crustacea, Invertebrata;

Bölüm II / Part II:

- (1) Uygun şekilde silin /
Delete as appropriate
(2) n = test edilen numune sayısı /
n= number of samples to be tested;

m = bakteri sayısı için eşik değeri; eğer tüm numunelerdeki bakteri sayısı bu değeri (m)'i aşmazsa sonuç olumlu değerlendirilir;/
m= threshold value for the number of bacteria: the result is considered satisfactory if the number of bacteria in all samples does not exceed m;

M = bakteri sayısı için maksimum değer; bir veya daha fazla numunedeki bakteri sayısı M veya daha fazla mikarda ise sonuç olumsuz olarak değerlendirilir; ve /
M= maximum value for the number of bacteria: the result is considered unsatisfactory if the number of bacteria in one or more samples is M or more:
and

C = bakteri sayısının "m" ile "M" arasında olabileceği numune sayısı, diğer numunelerdeki bakteri sayımı "m" veya bunun altında ise sonuç yine kabul edilebilir değerlendirilir. /
C= number of samples the bacterial count of which may be between m and M, the sample still being considered acceptable if the bacterial count of the other samples is m or less.

- İmza ve mühür renginin, baskından farklı bir renkte olması gerekmektedir /
The signature and the stamp must be in a different colour to that of the printing.
- Türkiye'de ki sevkiyattan sorumlu kişinin dikkatine: Bu sertifika yalnızca veteriner kullanım amaçlıdır ve ilgili sınır kontrol noktasına varana kadar sevkiyata eşlik etmelidir /
Note for the person responsible for the consignment in TR: this certificate is only for veterinary purposes and has to accompany the consignment until it reaches the border inspection post.

Bijlage 3: certificaat dogchews

**TÜRKİYE CUMHURİYETİ'NE KÖPEK ÇİĞNEME ÜRÜNLERİ İHRACATI İÇİN VETERİNER SAĞLIK SERTİFİKASI /
VETERINARY HEALTH CERTIFICATE FOR EXPORTATION OF DOGCHEWS TO THE REPUBLIC OF
TURKEY**

I.23. Konteynerin tanimlamasi/mühür numarası / Identification of container/seal number		I.24 Paketleme tipi / Type of packages			
1.25. Malların sertifikalandırma amacı / Commodities certified for: Pet hayvan yemi / <input checked="" type="checkbox"/> Petfood					
I.26. XXXXXXXXXXXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXXXXXXX	1.27. Türkiye'ye ithalat ya da kabul amaçlı / <input checked="" type="checkbox"/> For import into Turkey				
1.28. Malların tanımlaması / Identification of the commodities					
Parti numarası / Batch number	Türü (Bilimsel adı) / Species (Scientific name)	Malın tanımı / Nature of commodity	İşletmelerin Onay numarası: Üretim tesisi / Approval no of establishments: Manufacturing plant	Paket sayısı / Number of packages	Net ağırlık / Net weight

Ben, aşağıda imzası bulunan resmi veteriner hekim, Avrupa Parlamentosu ve Konseyi'nin 1069/2009/EC No'lu Yönetmeliği ve özellikle 10. Maddesini, Komisyon 142/2011/EU No'lu Yönetmeliği ve özellikle Ek XIII, Bölüm II ve Ek XIV, Bölüm II' sini okuduğumu ve anladığımı beyan eder ve yukarıda tanımlanan köpek çiğneme ürünlerine ilişkin olarak aşağıdakiler onaylarım:/ 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 and in particular Article 10 of that Regulation, and Commission Regulation (EU) No 142/2011, and in particular Chapter II of Annex XIII and Chapter II of Annex XIV thereto, and certify that the dogchews described above:

II.1. yalnızca aşağıdaki hayvansal yan ürünlerle hazırlanmıştır: / have been prepared exclusively with the following animal by- products:

ya⁽¹⁾ / either⁽¹⁾ İnsan tüketimine uygun olan, ancak ticari sebeplerle insan tüketimine sunulması amaçlanmayarak kesilen hayvanların karkas ve parçaları veya öldürülen av hayvanlarının gövdesi veya parçaları / carcasses and parts of animal slaughtered or, in the case of game, bodies or parts of animal killed, and which are fit for human consumption, but are not intended for human consumption for commercial reasons;

ve/veya⁽¹⁾ / and/or⁽¹⁾ bir kesimhanede kesilen ve antemortem bir muayeneyi takiben insan tüketimi için kesilmeye uygun bulunan hayvanlardan orjin alan karkaslar ve aşağıdaki parçaları veya insan tüketimi için öldürülen av hayvanlarının gövde ve aşağıdaki parçaları: / carcasses 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:

i) İnsan tüketimine uygun olmadığı için reddedilen, ancak insanlar veya hayvanlara geçebilen hastalık belirtisi göstermeyen hayvanların karkasları veya gövdeleri ile bunların parçaları / carcasses or bodies and parts of animals which are rejected as unfit for human consumption, but which did not show any signs of disease communicable to humans or animals;

ii) kanatlı kafaları / heads of poultry; hayvanların kirpintı ve parçaları dahil post ve derileri, boynuzları ve falanks, karpal ve merakarpal kemikleri, tarsal ve metatarsal kemikleri dahil ayakları /

- 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) domuz kilları / pig bristles;
 - v) tüyler / feathers;
- ve/veya⁽¹⁾ / and/or⁽¹⁾ antemortem bir muayeneyi takiben insan tüketimi için kesilmeye uygun bulunduktan sonra bir kesimhanede kesilmiş olan hayvanlardan elde edilen kan yolu ile insan veya hayvanlara bulaşabilecek herhangi bir hastalık belirtisi göstermeyen hayvanların kanları / blood of animals which did not show any signs of disease communicable through blood to humans or animals, obtained from animals that have been slaughtered in a slaughterhouse after having been considered fit for slaughter for human consumption following an ante- mortem inspection;
- ve/veya⁽¹⁾ / and/or⁽¹⁾ yağı alınmış kemikler, don yağı tortusu ve sütün işlenmesi sonucu ortaya çıkan santrifüj veya seperatör tortuları dahil insan tüketimi amaçlı ürünlerin üretiminden kaynaklanan hayvansal yan ürünler / animal by- products arising from the production of products intended for human consumption, including degreased bone, greaves and centrifuge or separator sludge from milk processing;
- ve/veya⁽¹⁾ / and/or⁽¹⁾ insan veya hayvanlara geçebilen herhangi bir hastalığın belirtisini göstermeyen, deniz memelileri hariç su hayvanları ve bu hayvanların parçaları / aquatic animals, and parts of such animals, except sea mammals, which did not show any signs of diseases communicable to humans or animals;
- ve/veya⁽¹⁾ / and/or⁽¹⁾ insan tüketimi için ürünler üreten işletme veya tesislerdeki su hayvanlarından elde edilen hayvansal yan ürünler / animal by-products from aquatic animals originating from plants or establishments manufacturing products for human consumption;
- ve/veya⁽¹⁾ / and/or⁽¹⁾ 96/22/EC nolu direktife istinaden yasaklı olan ve 1069/2009 No.lu Yönetmeliği (EC) Bölüm 35 (a) (ii)' ye uygun olarak malzemelerin ithaline izin verilen belirli maddeler ile muameleye tutulan hayvanlardan elde edilen malzemeler / material from animals which have been treated with certain substances which are prohibited pursuant to Directive 96/22/EC, the import of the material being permitted in accordance with Article 35(a)(ii) of Regulation (EC) No 1069/2009;
- II.2. Aşağıdaki işleme tabi tutulmuştur: / have been subjected:
- ya⁽¹⁾ / either⁽¹⁾ toynaklı hayvanların post ve derilerinden veya balıklardan elde edilmiş köpek çiğneme ürünleri olması durumunda, patojenik mikroorganizmaların (salmonella dahil) yok edilmesi için yeterli bir ısıl işleme tabi tutulmuştur ve bu köpek çiğneme ürünleri kurutulmuştur / in the case of dog chews made from hides and skins of ungulates or from fish, to a treatment sufficient to destroy pathogenic organisms (including salmonella); and the dog chews are dry;
- ve/veya⁽¹⁾ / and/or⁽¹⁾ toynaklı hayvanların post ve derileri veya balık dışındaki hayvansal yan ürünlerden elde edilmiş köpek çiğneme ürünleri olması durumunda, maddelerinin tamamı en az 90°C'lık bir ısıl işleme tabi tutulmuştur / in the case of dog chews made from animal by-products other than hides and skins of ungulates or from fish, to a heat treatment of at least 90° C throughout their substance;
- II.3. İşleme tesisinde depolama sırasında veya sonrasında her bir işlenmiş partiden alınan en az beş numune rastgele örneklemeye yoluyla incelenmiş ve aşağıdaki standartlara⁽²⁾ uygun bulunmuştur: / were examined by a random sampling of at least five samples from each processed batch taken during or after storage at the processing plant and complies with the following standards⁽²⁾:
- Salmonella : 25 g lik örnekte hiç yok: n=5, c=0, m=0, M=0 / absence in 25 g: n = 5, c = 0, m = 0, M = 0
- Enterobacteriaceae: 1 gr lik örnekte en çok n=5, c=2, m=10, M=300 / n = 5, c = 2, m = 10, M=300 in 1 gram;
- II.4. İşlem sonrasında patojenik ajanlarla kontaminasyonu engelleyecek tüm önlemler alınmıştır / have undergone all precautions to avoid contamination with pathogenic agents after treatment;
- II.5. Yeni ambalajında paketlenmiştir / were packed in new packaging;
- II.6. Yukarıda açıklanan köpek çiğneme ürünleri⁽¹⁾ / the dogchews described above⁽¹⁾

<p>ya⁽¹⁾ / either⁽¹⁾ veya⁽¹⁾ / or⁽¹⁾</p> <p>ya⁽¹⁾ / either⁽¹⁾ veya⁽¹⁾ / or⁽¹⁾</p>	<p>Sığır, koyun cinsi hayvanlar dışındaki ruminantlardan elde edilmişdir / is derived from other ruminants than bovine, ovine or caprine animals; Sığır, koyun ve keçi cinsi hayvanlardan elde edilmişdir ve aşağıdakileri içermez ve bunlardan türetilmemiştir / is derived from bovine, ovine or caprine animals and does not contain and is not derived from:</p> <p>Dünya Hayvan Sağlığı Teşkilatının (WOAH) sınıflandırmasına göre "ihmal edilebilir" BSE risk statüsünde yer alan bir ülkede veya bölgede doğmuş, sürekli olarak burada yetiştirilmiş ve kesilmiş hayvanlardan elde edilenler dışındaki sığır, koyun veya keçi cinsi hayvanların ürünlerini / bovine, ovine or 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 WOAH;</p> <p>(a) Avrupa Konseyi ve Parlamentosu'nun 999/2001 (EC) sayılı Yönetmeliği'nin Ek V' in 1. maddesinde tanımlanan spesifik risk materyali / specified risk material as defined in point 1 of Annex V to Regulation (EC) No 999/2001 of the European Parliament and of the Council;</p> <p>(b) Yöresel olarak BSE vakası bulunmayan, Dünya Hayvan Sağlığı Teşkilatı (WOAH) standartlarına göre ihmal edilebilir BSE risk statüsünde sınıflandırılmış bir ülkede veya bölgede sürekli olarak yetiştirilen ve kesilen sığır, koyun, keçi cinsi hayvanlar dışındaki hayvanların kemiklerinden mekanik olarak ayrılmış olan etleri / mechanically separated meat obtained from bones of bovine, ovine or caprine animals, except from those animals that were born, continuously reared and slaughtered in a country or region classified as posing a negligible BSE risk in accordance with WOAH, in which there has been no indigenous BSE case;</p> <p>(c) Dünya Hayvan Sağlığı Teşkilatı (WOAH) standartlarına göre ihmal edilebilir BSE risk statüsündeki ülke veya bölgede doğmuş, sürekli yetiştirilmiş ve kesilmiş hayvanlar dışında kafatası boşluğununa sokulmuş uzun çubuk şeklinde bir alet vasıtasyyla merkezi sinir sistemi dokularının harap edilmesi yoluyla veya kafatası boşluğununa gaz enjeksiyonu yoluyla sersemletme sonrasında öldürülün sığır, koyun veya keçi cinsi hayvanlardan elde edilen hayvansal yan ürün ve türev ürünleri / 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 WOAH.</p>
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Notlar / Notes

Bölüm I / Part I:

- Kutu referansı I.15: Sicil numarası (tren vagonları veya konteynır ya da kamyonlar), uçuş
sayısı (uçak) veya isim (gemi); bilgiler boşaltma veya yeniden yükleme halinde verilmelidir /
Box reference 1.15: Registration number (railway wagons or container and lorries), flight
number (aircraft) or name (ship); information is to be provided in the event of unloading
and reloading;
- Kutu referansı I.19: 05.11, 23.09, 41.01 veya 4205 /
Box reference 1.19: 05.11, 23.09, 41.01 or 4205.
- Kutu referansı I.23: Dökme yük konteynırlar için, konteynır numarası ve mühür numarası
(varsayı) verilmelidir./
Box reference I.23: for bulk containers, the container number and the seal number (if
applicable) should be given.
- Kutu referansı I.25: teknik kullanım: Çiftlik hayvanlarının, kürk hayvanlarının dışındaki
hayvanların beslenmesi ve pet hayvan yemlerinin üretimi ya da imalatı için kullanılması /

Box reference I.25: technical use: any use other than for feeding of farmed animals, other than fur animals, and the production or manufacturing of pet food.

Bölüm II / Part II:

- (1) Uygun şekilde silin /
Delete as appropriate
- (2) Where:
 n = test edilen numune sayısı /
 n = number of samples to be tested;

m = bakteri sayısı için eşik değeri; eğer tüm numunelerdeki bakteri sayısı bu değeri (m)'ı aşmazsa sonuç olumlu değerlendirilir;/
 m = threshold value for the number of bacteria: the result is considered satisfactory if the number of bacteria in all samples does not exceed m ;

M = bakteri sayısı için maksimum değer; bir veya daha fazla numunedeki bakteri sayısı M veya daha fazla miktarda ise sonuç olumsuz olarak değerlendirilir; ve /
 M = maximum value for the number of bacteria: the result is considered unsatisfactory if the number of bacteria in one or more samples is M or more: and

C = bakteri sayısının " m " ile " M " arasında olabileceği numune sayısı, diğer numunelerdeki bakteri sayımı " m " veya bunun altında ise sonuç yine kabul edilebilir değerlendirilir. /
 C = number of samples the bacterial count of which may be between m and M , the sample still being considered acceptable if the bacterial count of the other samples is m or less.

- İmza ve mühür renginin, baskından farklı bir renkte olması gerekmektedir /
The signature and the stamp must be in a different colour to that of the printing
- Türkiye'de ki sevkiyattan sorumlu kişinin dikkatine: Bu sertifika yalnızca veteriner kullanım amaçlıdır ve ilgili sınır kontrol noktasına varana kadar sevkiyata eşlik etmelidir /
Note for the person responsible for the consignment in TR: this certificate is only for veterinary purposes and has to accompany the consignment until it reaches the border inspection post