## UNITED STATES Official certificate to the EU

### Part I: Description of consignment

<table>
<thead>
<tr>
<th>Description</th>
<th>Code</th>
</tr>
</thead>
<tbody>
<tr>
<td>I.1 Consignor/Exporter Name</td>
<td></td>
</tr>
<tr>
<td>Address</td>
<td></td>
</tr>
<tr>
<td>Country ISO country code</td>
<td></td>
</tr>
<tr>
<td>I.5 Consignee/Importer Name</td>
<td></td>
</tr>
<tr>
<td>Address</td>
<td></td>
</tr>
<tr>
<td>Country ISO country code</td>
<td></td>
</tr>
<tr>
<td>I.7 Country of origin ISO country code</td>
<td></td>
</tr>
<tr>
<td>I.8 Region of origin Code</td>
<td></td>
</tr>
<tr>
<td>I.11 Place of dispatch Name</td>
<td></td>
</tr>
<tr>
<td>Registration/Approval No</td>
<td></td>
</tr>
<tr>
<td>Address</td>
<td></td>
</tr>
<tr>
<td>Country ISO country code</td>
<td></td>
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</tbody>
</table>

### Description of consignment

<table>
<thead>
<tr>
<th>Description</th>
<th>Code</th>
</tr>
</thead>
<tbody>
<tr>
<td>CN code</td>
<td>Cold Store</td>
</tr>
<tr>
<td>Slaughterhouse</td>
<td>Treatment type</td>
</tr>
</tbody>
</table>

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Signature of Official Veterinarian or Official Inspector
Country: United States

Certificate Edition (05/2022)  
FSIS Form 2630-9 (6/86)  

II. Health information

I. the undersigned, hereby certify that


II.2. the composite products described in Part I:

(a) comply with Article 5 of Regulation (EC) No 852/2004, in particular they come from (an) establishment(s) implementing a programme based on the hazard analysis and critical control points (HACCP) principles, regularly audited by the competent authorities;

(b) comply with Article 6(1), point (b), of Regulation (EC) No 853/2004 on the origin of the products of animal origin used in their production;

(c) were produced in accordance with the requirements referred to under point II.1.;

(d) fulfil the guarantees covering live animals and products thereof provided by the residue plans submitted in accordance with Article 29 of Council Directive 96/23/EC;

(e) contain processed products of animal origin that where produced in establishments located in European Union Member States or in third countries authorised for entry into the European Union of those processed products of animal origin;

(f) have been produced under conditions guaranteeing compliance with the maximum residue levels for pesticides laid down in Regulation (EC) No 396/2005, and the maximum levels for contaminants laid down in Regulation (EC) No 1881/2006.

II.3. the composite products described in Part I contain:

(1) [II.3.A. Meat products(2) in any quantity except gelatine, collagen and highly refined products referred to in Annex III, Section XVI, to Regulation (EC) No 853/2004, which:

1) meet the animal health requirements laid down in Commission Delegated Regulation (EU) 2020/692 and contain the following meat constituents which are eligible for entry into the Union as such and meet the following criteria:

<table>
<thead>
<tr>
<th>Species (3)</th>
<th>Treatment (4)</th>
<th>Origin (5)</th>
<th>Approved Establishment(s) (6)</th>
</tr>
</thead>
</table>

(1) [2] originate from

(1) either [the same country as the country of origin in Box I.7;]

(1) and/or [a Member State;]

(1) and/or [a third country or parts thereof authorised for entry into the Union of meat products not required to undergo a specific risk-mitigating treatment as set out in Annex XV to Commission Implementing Regulation (EU) 2021/404, and the third country where the composite product is produced is also authorised for entry into the Union of meat products treated with that treatment.]

(1) [3] if containing material from bovine, ovine or caprine animals, with regard to bovine spongiform encephalopathy (BSE):
**II. Health information**

(1) either [the country or region of origin is classified in accordance with Commission Decision 2007/453/EC as a country or region posing a negligible BSE risk, and

(1) [the animals from which the meat products are derived were born, continuously reared and slaughtered in a country or region classified in accordance with Decision 2007/453/EC as a country or region posing a negligible BSE risk in which there have been no BSE indigenous cases;]

(1) [the animals from which the meat products are derived originate from a country or region classified in accordance with Decision 2007/453/EC as a country or region posing a negligible BSE risk in which there has been at least one BSE indigenous case, and the meat products do not contain and are not derived from mechanically separated meat obtained from bones of bovine, ovine and caprine animals;]

(1) [the animals from which the meat products are derived originate from a country or region classified in accordance with Decision 2007/453/EC as a country or region posing a controlled BSE risk and:

(i) the meat products do not contain and are not derived from specified risk material as defined in Annex V, point 1, to Regulation (EC) No 999/2001 of the European Parliament and of the Council;

(ii) the meat products do not contain and are not derived from mechanically separated meat obtained from bones of bovine, ovine and caprine animals;

(iii) the animals from which the meat products are derived have not been slaughtered after stunning by means of gas injected into the cranial cavity or killed by the same method or slaughtered by laceration after stunning of central nervous tissue by means of an elongated rod-shaped instrument introduced into the cranial cavity;]

(1) [the animals from which the meat products are derived originate from a country or region classified in accordance with Decision 2007/453/EC as a country or region posing an undetermined BSE risk and:

(i) the meat products do not contain and are not derived from specified risk material as defined in Annex V, point 1, to Regulation (EC) No 999/2001;

(ii) the meat products do not contain and are not derived from mechanically separated meat obtained from bones of bovine, ovine and caprine animals;

(iii) the animals from which the meat products are derived have not been slaughtered after stunning by means of gas injected into the cranial cavity or killed by the same method or slaughtered by laceration after stunning of central nervous tissue by means of an elongated rod-shaped instrument introduced into the cranial cavity;

(iv) the animals from which the meat products are derived have not been fed with meat-and-bone meal or greaves, as defined in the Terrestrial Animal Health Code of the World Organisation for Animal Health;

(v) the meat products were produced and handled in a manner which ensures that they do not contain and were not contaminated with nervous and lymphatic tissues exposed during the deboning process;]
II. Health information

(1) and/or [the country or region of origin is classified in accordance with Decision 2007/453/EC as a country or region posing a controlled BSE risk, and

(a) the animals from which the meat products are derived have not been slaughtered after stunning by means of gas injected into the cranial cavity or killed by the same method or slaughtered by laceration after stunning of central nervous tissue by means of an elongated rod-shaped instrument introduced into the cranial cavity;

(1) or [the meat products do not contain and are not derived from:

(i) specified risk material as defined in Annex V, point 1, to Regulation (EC) No 999/2001;

(ii) mechanically separated meat obtained from bones of bovine, ovine and caprine animals;]

(1) or [the meat products contain and are derived from treated intestines sourced from animals which were born, continuously reared and slaughtered in a country or region classified in accordance with Decision 2007/453/EC as a country or region posing a negligible BSE risk in which there have been no BSE indigenous cases;]

(1) or [the meat products contain and are derived from treated intestines sourced from animals which originate from a country or region classified in accordance with Decision 2007/453/EC as a country or region posing a negligible BSE risk in which there has been at least one BSE indigenous case, and:

(1) either [(i) the animals were born after the date from which the ban on the feeding of ruminants with meat-and-bone meal and greaves derived from ruminants has been enforced;]

(1) or [(ii) the treated intestines of bovine, ovine and caprine animal origin do not contain and are not derived from specified risk material as defined in Annex V, point 1, to Regulation (EC) No 999/2001;]

(1) either [(c) the animals from which the meat products are derived originate from a country or region classified in accordance with Decision 2007/453/EC as a country or region posing a negligible or a controlled BSE risk;]

(1) or [(c) the animals from which the meat products are derived originate from a country or region classified in accordance with Decision 2007/453/EC as a country or region posing an undetermined BSE risk and

(i) the animals from which the meat products are derived have not been fed with meat-and-bone meal or greaves, as defined in the Terrestrial Animal Health Code of the World Organisation for Animal Health;

(ii) the meat products were produced and handled in a manner which ensures that they do not contain and were not contaminated with nervous and lymphatic tissues exposed during the deboning process;]

(1) and/or [the country or region of origin is classified in accordance with Decision 2007/453/EC as a country or region with an undetermined BSE risk, and

(a) the animals from which the meat products are derived have not been:

(i) slaughtered after stunning by means of gas injected into the cranial cavity or killed by the same method or slaughtered by laceration after stunning of central nervous tissue by means of an elongated rod-shaped instrument introduced into the cranial cavity;]
II. Health information

(ii) fed meat-and-bone meal or greaves derived from ruminants, as defined in the Terrestrial Animal Health Code of the World Organisation for Animal Health;

(1) either (b)

the meat products do not contain and are not derived from:

(i) specified risk material as defined in Annex V, point 1, to Regulation (EC) No 999/2001;

(ii) mechanically separated meat obtained from bones of bovine, ovine and caprine animals;

(iii) nervous and lymphatic tissues exposed during the deboning process.

(1) or (b)

the meat products contain and are derived from treated intestines sourced from animals which were born, continuously reared and slaughtered in a country or region classified in accordance with Decision 2007/453/EC as a country or region posing a negligible BSE risk in which there have been no BSE indigenous cases;

(1) or

the meat products contain and are derived from treated intestines sourced from animals which originate from a country or region classified in accordance with Decision 2007/453/EC as a country or region posing a negligible BSE risk in which there has been at least one BSE indigenous case, and:

(1) either (i)

the animals were born after the date from which the ban on the feeding of ruminants with meat-and-bone meal and greaves derived from ruminants has been enforced;

(1) or (i)

the treated intestines of bovine, ovine and caprine animal origin do not contain and are not derived from specified risk material as defined in Annex V, point 1, to Regulation (EC) No 999/2001.

II.3.B. Dairy products or colostrum-based products(8) in any quantity that meet the animal health and/or requirements laid down in Commission Delegated Regulation (EU) 2020/692 and therefore are eligible for entry into the Union as such, and:

(a) have been produced

(1) either [in the zone with code __________ as listed in Annex XVII, Part 1, to Implementing Regulation (EU) 2021/404 which has been free from foot and mouth disease and infection with rinderpest virus for the period of at least 12 months prior to the date of milking and, during that period, no vaccination against those diseases has been carried out.]

(1) and/or [in the zone with code __________ as listed in Annex XVIII, Part 1, to Implementing Regulation (EU) 2021/404 and the treatment applied complies with the minimum treatment provided for in Article 157 and Annex XXVII to Delegated Regulation (EU) 2020/692]

(1) and/or [in a Member State;]

and in the establishment __________ (approval number of the establishments of origin of the dairy products or the colostrum-based products contained in the composite product authorised at the time of production for export of dairy products or colostrum-based products to the European Union).

(b) originate in:

(1) either [the same zone as the zone referred to in Box I.7.]

(1) and/or [a Member State;]
### II. Health information

| (1) | and/or [a zone authorised for entry into the Union of milk, colostrum, dairy products and colostrum-based products in Annex XVII, Part 1, to Implementing Regulation (EU) 2021/404, where the zone where the composite product is produced is also authorised, under the same conditions, for entry into the Union of milk, colostrum, dairy products and colostrum-based products and listed in Part 1 of that Annex;] |
| (1) | (c) are dairy products made from raw milk obtained from [Bos Taurus](1), [Ovis aries](1), [Capra hircus](1), [Bubalus bubalis](1), [Camelus dromedarius](1) and prior to dispatch to the Union have undergone or been produced from raw milk which has undergone: |
| (1) | either [a pasteurisation treatment involving a single heat treatment with a heating effect at least equivalent to that achieved by a pasteurisation process of at least 72°C for 15 seconds and where applicable, sufficient to ensure a negative reaction to an alkaline phosphatase test applied immediately after the heat treatment;] |
| (1) | or [a sterilisation process, to achieve an F0 value equal to or greater than three;] |
| (1) | or [an ultra high temperature (UHT) treatment at not less than 135°C in combination with a suitable holding time;] |
| (1) | or [a high temperature short time pasteurisation treatment (HTST) at 72°C for 15 seconds, or a treatment with an equivalent pasteurisation effect, applied to milk with a pH lower than 7,0 achieving, where applicable, a negative reaction to an alkaline phosphatase test;] |
| (1) | or [a high temperature short time pasteurisation treatment (HTST) at 72°C for 15 seconds, or a treatment with an equivalent pasteurisation effect, applied twice to milk with a pH equal to or greater than 7,0 achieving, where applicable, a negative reaction to an alkaline phosphatase test, immediately followed by] |
| (1) | either [lowering the pH below 6 for one hour;] |
| (1) | or [additional heating equal to or greater than 72°C, combined with desiccation;] |
| (1) | or animals other than Bos Taurus, Ovis aries, Capra hircus, Bubalus bubalis and Camelus dromedarius and prior to dispatch to the Union have undergone or been produced from raw milk which has undergone |
| (1) | either [a sterilisation process, to achieve an F0 value equal to or greater than three;] |
| (1) | or [an ultra high temperature (UHT) treatment at not less than 135°C in combination with a suitable holding time;] |
| (1) | (d) are colostrum-based products and they come from a third country or territory listed in Annex XVII, Part 1, to Implementing Regulation (EU) 2021/404 for entry into the Union of raw milk, colostrum and colostrum-based products |
| (1) | (e) were produced on or between and [9).] |
| (1) | (II.3.C.) Fishery products that originate from the approved establishment N°(10) |
| (1) | situated in the country(11) ] |
| (1) | (II.3.D.1.) originate from |
| (1) | (II.3.D.) Egg products that |
| (1) | (I) and/or [the zone(12) which at the date of issue of this animal health/official certificate is listed in Annex XIX, Part 1 to Implementing Regulation (EU) 2021/404 for the entry into the Union of egg products and applies a disease surveillance programme for highly pathogenic avian influenza that complies with the requirements referred to in Article 160 of Delegated Regulation (EU) 2020/692;] |
II. Health information

(1) and/or [a Member State;]

II.3.D.2. were produced from eggs coming from an establishment which satisfies the requirements of Annex III, Section X, to Regulation (EC) No 853/2004 in which, during the period of at least 30 days prior to the date of collection of the eggs, no outbreak of highly pathogenic avian influenza and infection with Newcastle disease virus has occurred and:

(1) either [(a) within a 10 km radius of which, including, where appropriate, the territory of a neighbouring country, there has been no outbreak of highly pathogenic avian influenza during the period of at least 30 days prior to the date of the collection of the eggs;]

(1) or [(a) the egg products have undergone the following treatment:]

(1) either [liquid egg white was treated:
(1) either [with 55.6°C for 870 seconds;]
(1) or [with 56.7°C for 232 seconds;]
(1) or [10% salted yolk was treated with 62.2°C for 138 seconds;]
(1) or [dried egg white was treated:
(1) either [with 67°C for 20 hours;]
(1) or [with 54.4°C for 50.4 hours;]
(1) or [whole eggs were:
(1) either [at least treated with 60°C for 188 seconds;]
(1) or [completely cooked;]
(1) or [whole egg blends were at least treated:
(1) either [with 60°C for 188 seconds;]
(1) or [with 61.1°C for 94 seconds;]
(1) or [completely cooked;]
(1) either [within a 10 km radius of which, including where appropriate, the territory of a neighbouring country there was no outbreak of infection with Newcastle disease virus during the period of at least 30 days prior to the date of collection of the eggs;]

(1) or [(b) the egg products have undergone the following treatment:]

(1) either [liquid egg white was treated:
(1) either [with 55°C for 2278 seconds;]
(1) or [with 57°C for 986 seconds;]
(1) or [with 59°C for 301 seconds;]
(1) or [10% salted yolk was treated with 55°C for 176 seconds;]
(1) or [dried egg white was treated with 57°C for 50.4 hours;]
(1) or [whole eggs were:
(1) either [treated with 55°C for 2521 seconds;]
(1) or [treated with 57°C for 1596 seconds;]
(1) or [treated with 59°C for 674 seconds;]
(1) or [completely cooked;]]]
Country: United States

II. Health information

Notes

In accordance with the Agreement on the withdrawal of the United Kingdom of Great Britain and Northern Ireland from the European Union and the European Atomic Energy Community, and in particular Article 5(4) of the Protocol on Ireland / Northern Ireland in conjunction with Annex 2 to that Protocol, references to European Union in this animal health/official certificate include the United Kingdom in respect of Northern Ireland.

This animal health/official certificate shall be completed in accordance with the notes for the completion of certificates provided for in Annex I, Chapter 4, to Implementing Regulation (EU) 2020/2235.

Part I:


Box  Name, address and registration/approval number if available of the establishments of production of the composite product(s). Name of the country of dispatch which must be the same as the country of origin in Box I.7.

Box  Registration number (railway wagons or container and road vehicles), flight number (aircraft) or name (vessel). In the case of transport in containers their registration number and where there is a serial number of the seal it must be indicated in Box I.19. In the case of unloading and reloading, the consignor must inform the border control post of entry into the Union.

Box  For containers or boxes, the container number and the seal number (if applicable) must be included.

Box  Use the appropriate Harmonised System (HS) code of the World Customs Organisation such as: 1517, 1518, 1601 00, 1602, 1603 00, 1604, 1605, 1702, 1704, 1806, 1901, 1902, 1904, 1905, 2001, 2004, 2005, 2101, 2103, 2104, 2105 00, 2106, 2202, 2208.

Box  Description of consignment:

“Manufacturing plant”:

Insert the name and approval number if available of the establishments of production of the composite product(s).

“Nature of commodity”:

In the case of composite products containing meat products indicate ‘meat product’. In the case of composite product containing dairy products indicate ‘dairy product’. In the case of composite product containing colostrum-based products indicate ‘colostrum-based product’. In the case of composite product containing fishery products specify whether aquaculture or wild origin. In the case of composite product containing egg products indicate ‘egg products’.

Part II:

(1)  Keep as appropriate.


(3)  Insert the code for the relevant species of the meat product where BOV = domestic bovine animals (Bos taurus, Bison bison, Bubalus bubalis and their crossbreds); OVI = domestic sheep (Ovis aries) and goats (Capra hircus); EQU = domestic equine animals (Equus caballus, Equus asinus and their crossbreds), POR = domestic porcine animals (Sus scrofa); RM = farmed rabbits, POU = domestic poultry, RAT = ratites, RUF: animals of the family Bovidae (other than domestic bovine, ovine and caprine animals), camelid animals and cervid animals kept as farmed game; RUW: wild animals of the family Bovidae (other than domestic bovine, ovine and caprine animals), wild camelid animals and wild cervid animals; SUF: animals kept as farmed game of wild breeds of porcine animals and animals of the family Tayassuidae; SUW: wild animals of wild breeds of porcine animals and animals of the family Tayassuidae; EQW = wild game solipeds, WL = wild leporidae, WM = wild land mammals other than ungulates and leporidae; GBM = Game birds.
Country: United States

<table>
<thead>
<tr>
<th>Part II: Certification</th>
<th>Certificate reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>(4) Insert A, B, C, D, E or F for the required treatment as specified and defined in Annex XV to Implementing Regulation (EU) 2021/404.</td>
<td></td>
</tr>
<tr>
<td>(5) Insert the code of the zone of origin of the meat product, as listed in Annex XV to Implementing Regulation (EU) 2021/404.</td>
<td></td>
</tr>
<tr>
<td>(6) Insert EU approval number of the establishments of origin of the meat products contained in the composite product.</td>
<td></td>
</tr>
<tr>
<td>(7) Delete if the meat products are obtained from EQU, EQW, WL, RM or WM or as defined in footnote (3).</td>
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</tr>
<tr>
<td>(9) Date or dates of production. Composite products shall only be permitted to enter into the Union if the products of animal origin contained therein were obtained after the date of authorisation of the third country or part thereof where the products of animal origin were produced, for entry into the Union of the specific species and category of products of animal origin, or during a period where animal health restriction measures taken by the European Union were not in place against the entry of those products from this third country or part thereof, or during a period where the authorisation of this country or part thereof for entry into the Union of those products was not suspended.</td>
<td></td>
</tr>
<tr>
<td>(10) Number of the fishery product establishment authorised to export to the European Union.</td>
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</tr>
<tr>
<td>(11) Country of origin authorised for entry into the Union. In the case of fishery products derived from bivalve molluscs the country of origin must be authorised for entry into the Union of live bivalve molluscs.</td>
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<tr>
<td>(13) to be signed by:</td>
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<tr>
<td>- an official veterinarian,</td>
<td></td>
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<tr>
<td>- a certifying officer or an official veterinarian for composite products containing only egg or fishery products.</td>
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<tr>
<td>(14) Keep at least one of the proposed options.</td>
<td></td>
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</tbody>
</table>

Official veterinarian or Official inspector

<table>
<thead>
<tr>
<th>Name (in capital letters)</th>
<th>Qualification and title</th>
</tr>
</thead>
<tbody>
<tr>
<td>Date of signature</td>
<td>Signature</td>
</tr>
<tr>
<td>Stamp</td>
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