| | I.1. Consignor | | | I.2. IMSOC Reference | I.2. IMSOC Reference | | |
|---------------------------------|---|-------------------------|----------------------------------|---|---|--|--|
| | Name | | | I.2.a. Local Reference | | | |
| | Address | | | | | | |
| | Country | | ISO Code | | | | |
| | I.5. Consignee | | | I.3. Central competent authority | | | |
| ┙ | | | | | | | |
| Part I : Details of consignment | Address | | | I.4. Local competent authority | | | |
| | Country | | ISO Code | | | | |
| 뛆 | , | | | | | | |
| NS | I.7. Country of ori | gin | ISO Code | I.9. Country of destination | ISO Code | | |
| ខ | | | | | | | |
| 비 | I.8. Region of origi | in | Code | I.10. Region of destination | | | |
| ils | I.11. Place of Dispa | atch | | I.12. Place of destination | | | |
| ŝta | Name | | | Name | | | |
| Ă | Address | | | Address | | | |
| н | Approval Numbe | r | | Approval Number | | | |
| 빏 | Country | | ISO Code | Country | ISO Code | | |
| Ä | I.13. Place of Load | ing | | I.14. Date and time of departure | | | |
| | Name | | | | | | |
| | Address | | | | | | |
| | Approval Numbe | r | | | | | |
| | Country | | ISO Code | | | | |
| | - | | | | | | |
| _ | I.15. Means of Tra | | x 1 | I.16 Entry Point | | | |
| | Mode | International transport | Identification | | | | |
| | | document | | | | | |
| | | | | | | | |
| | | | | | | | |
| | | | | | | | |
| | | | | | | | |
| | I.18. Transport co | nditions | | I.17. Accompanying documents | | | |
| | Ambient 🗆 | Chilled 🗆 | Controlled Frozen 🗆 | Accompanying document | | | |
| | | | temperature | reference | | | |
| | | | | Date of issue | | | |
| | | | | Country Place of issue | | | |
| | I.19. Container No | / Seal No | | | | | |
| | | | | | | | |
| | I.20. Certified as | | | | | | |
| | Breeding 🗆 | | Relaying 🗆 | Artificial reproduction \Box | Fattening 🗆 | | |
| | Animal Feedingstuff 🗆 Pharmaceutical use 🗖 | | Technical use 🗖 | Other 🗆 | Human consumption \square | | |
| | | | Slaughter 🗆 | Production of petfood \Box | Breeding and production \Box | | |
| | Production 🗆 | | 0 | - | Ŭ I | | |
| | | | | | | | |
| | I.21. For transit th | rough a third cour | ntry | I.22. For transit through Member | State(s) | | |
| | Country | | ISO Code | Country | ISO Code | | |
| | EU Exit | | BCP code | | | | |
| | Authority | | | | | | |
| | EU Entry Authority | | BCP code | | | | |
| | I.23. Total number | r of packages | I.24. Total quantity | I.25. Total net weight | I.25. Total gross weight | | |
| | | | | | | | |
| | I.28. Description o | - | | | | | |
| | 1. 35 ALBUMINOI | DAL SUBSTANCES | ; MODIFIED STARCHES; GLUES; E | NZYMES | | | |
| | 3504 Peptones a | and their derivativ | es; other protein substances and | their derivatives, not elsewhere specifie | ed or included; hide powder, whether or | | |
| | not chrômed #1. Commodity | | Quantity | Net weight | Package count | | |
| | | | | | | | |
| | Species | | Identification number | Identification system | | | |
| | | | | | | | |
| | | | | | | | |
| | | | | | | | |
| | | | | | | | |
| | | | | | | | |
| | | | | | | | |

| | II. Health infor | rmation | | | | | | |
|------------------------|---|---|---|--|------------------------------|--|--|--|
| Part II: Certification | 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 (c) and Article 8(d) and Article 10 thereof, and Commission Regulation (EU) No 142/2011, and in particular Chapter II of Annex XIV thereto, and certify that: | | | | | | | |
| | II.1. | the blood p | roducts described above consist of | plood products that satisfy the requirements below; | | | | |
| | II.2. | they consis | t exclusively of blood products not | ntended for human or animal consumption; | | | | |
| | II.3. | - | been prepared and stored in a plant ng animal by-products: | supervised by the competent authority, exclusively with | | | | |
| Certif | (2) | 🗆 either | [- blood of slaughtered animals, which is fit for human consumption in accordance with retained EU law, but is not intended for human consumption for commercial reasons;] | | | | | |
| Part II: | (2) | □ and/or | accordance with retained EU law, l communicable to humans or anim a slaughterhouse and were conside | of slaughtered animals, which is rejected as unfit for human consumption in the with retained EU law, but which did not show any signs of diseases icable to humans or animals, derived from carcasses that have been slaughtered in terhouse and were considered fit for human consumption following an ante- inspection in accordance with retained EU law;] | | | | |
| | (2) | □ and/or | [- blood of slaughtered animals, which did not show any signs of diseases communicable to humans or animals, obtained from animals that have been slaughtered in a slaughterhouse after having been considered fit for human consumption following an ante-mortem inspection in accordance with retained EU law;] | | | | | |
| | (2) | □ and/or [- blood and blood products originating from live animals that did not show clinical signs of any disease communicable through these products to humans or animals;] | | | | | | |
| | (2) | □ and/or | [- blood and blood products derive consumption;] | ed from the production of products intended for human | | | | |
| | (2) | □ and/or | [- animal by-products which have l illegal treatment as defined in Arti Council Directive 96/23/EC;] | | | | | |
| | (2) | □ and/or | contaminants listed in Group B(3) | residues of other substances and environmental of Annex I to Directive 96/23/EC, if such residues exceed retained EU law or, in the absence thereof, in national | | | | |
| | II.4. | the blood that these products were manufactured from has been collected in slaughterhouses approved in accordance with retained EU law, in slaughterhouses approved and supervised by the competent authority of the country of collection or from live animals in facilities approved and supervised by the competent authority of the country of collection. | | | | | | |
| | (2) 🗆 [II.5. | In the case of blood products derived from Artiodactyla, Perissodactyla and Proboscidea including their crossbreeds, other than Suidae and Tayassuidae, the products have undergone one of the following treatments, guaranteeing the absence of pathogens of foot-and-mouth disease, vesicular stomatitis, rinderpest, peste des petits ruminants, Rift Valley fever and bluetongue: | | | | | | |
| | (2) | 🗆 either | [heat treatment at a temperature of 65°C for at least three hours, followed by an effectiveness check;] | | | | | |
| | (2) | \Box and/or | [irradiation at 25 kGy by gamma ra | ays, followed by an effectiven | ess check;] | | | |
| | (2) | \Box and/or | [change in pH to pH 5 for two hour | rs, followed by an effectivene | ss check;] | | | |
| | (2) | □ and/or | [heat treatment of at least 80 °C the check]] | oughout their substance, foll | owed by an effectiveness | | | |
| | (2) 🗆 [II.6. | products ha following d | of blood products derived from Suidae, Tayassuidae, poultry and other avian species, the ave undergone one of the following treatments guaranteeing the absence of pathogens of the iseases: foot-and-mouth disease, vesicular stomatitis, swine vesicular disease, classical swine an swine fever, Newcastle disease and highly pathogenic avian influenza, as appropriate to | | | | | |
| | (2) | 🗆 either | [heat treatment at a temperature o effectiveness check;] | f 65°C for at least three hours, followed by an | | | | |
| | (2) | \Box and/or | [irradiation at 25 kGy by gamma ra | ays, followed by an effectiven | ess check;] | | | |
| | (2) | □ and/or | [heat treatment of at least 80°C for | Suidae/Tayassuidae (2) and a | t least 70°C for poultry and | | | |

| | II. Health info | rmation | | | | | | | |
|------------------------|------------------------|--|--|-------------|---|--|---|--|--|
| | | other avian species (2) throughout the substance of the product, followed by an effectiveness check]]. | | | | | | | |
| Part II: Certification | (2) 🗆 [II.7. | | e of blood products derived from species other than those listed in point II.5 or II.6, the nave undergone the following treatment (please specify): .] | | | | | | |
| | II.8. | The produ | icts were: | | | | | | |
| | (2) | \circ either | [packed in new or sterilised bags or bottles,] | | | | | | |
| | (2) | ∘ or | [transported in bulk in containers or other means of transport that were thoroughly cleaned and disinfected with a disinfectant approved by the competent authority before use;] | | | | | | |
| | | | and the outer packaging or containers bear labels indicating 'NOT FOR HUMAN OR ANIMAL CONSUMPTION'; | | | | | | |
| | II.9. | the produc | cts were stored in enclosed storage; | | | | | | |
| | II.10. | all precaut treatment; | ll precautions were taken to avoid the contamination of the products with pathogenic agents after reatment; | | | | | | |
| | (2) 🗆 [II.11. | The treated | ated blood products described above | | | | | | |
| | (2) | \circ either | [is derived from other ruminants than bovine, ovine or caprine animals.]] | | | | | | |
| | (2) | ∘ or | [is derived from bovine, ovine or caprine animals and does not contain and is not derived from: | | | | | | |
| | born, continuously rea | | | nuously rea | orine materials other than those derived from animals ared and slaughtered in a country or region classified as E risk in accordance with Decision 2007/453/EC.]] | | | | |
| | | (2) | o or | [(a) | - | sk material as defined in po 9/2001 of the European Parli | int 1 of Annex V to Regulation ament and of the Council; | | |
| | | | | (b) | or caprine continuous classified a | animals, except from those ly reared and slaughtered i s posing a negligible BSE ris n Decision 2007/453/EC, wh | n a country or region k in accordance with | | |
| | | | | (c) | or caprine laceration of rod-shaped means of g animals that country or | animals which have been k of the central nervous tissue i instrument introduced into as injected into the cranial o | by means of an elongated the cranial cavity, or by avity, except for those reared and slaughtered in a negligible BSE risk in | | |
| | | | | | | | | | |

Notes

(*) Those countries subject to the transitional import arrangements include: an EU member State; Liechtenstein; Norway; Iceland and Switzerland.

References to European Union legislation within this certificate are references to direct EU legislation which has been retained in Great Britain (retained EU law as defined in the European Union (Withdrawal) Act 2018).

References to Great Britain in this certificate include Channel Islands and Isle of Man.

Part I:

BoxPerson responsible for the consignment in Great Britain, Channel Islands or Isle of Man: this
referenceI.6:Box is required to be filled in only if it is a certificate for a commodity to be transited
through Great Britain, Channel Islands or Isle of Man; it may be filled in if the certificate is
for a commodity to be imported into Great Britain, Channel Islands or Isle of Man.BoxApproval number; the registration number of the establishment or plant, which has been
issued by the competent authority.

| | | | (, | | | | |
|---|--|--|---|--------------------------|--|--|--|
| II. Health ini | formation | | | | | | |
| | I.11 and I.12: | | | | | | |
| - | Box reference I.12: | Place of destination; this box is to be filled in only if it is a certificate for a transit commodity. Products in transit may only be stored in free zones, free warehouses and custom warehouses. | | | | | |
| | Box reference I.15: | Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In the case of unloading and reloading in Great Britain, Channel Islands or Isle of Man, the consignor must inform the BCP of entry into Great Britain, Channel Islands or Isle of Man. | | | | | |
| | Box reference I.16: | Do not use this box until the end of the transitional staging period. | | | | | |
| - | Box reference I.19: | use the appropriate Harmonized System (HS) code under the following headings: 05.11, 30.02, 35.02 or 35.04. | | | | | |
| - | Box reference I.23: | for bulk containers, the container number and the seal number (if applicable) must be included. | | | | | |
| - | Box reference I.25: | technical use: any use other than f production or manufacturing of po | er than feeding of farmed animals, other than fur animals, and the ing of pet food. | | | | |
| - | Box reference I.26 and I.27: | fill in according to whether it is a transit or an import certificate. e | | | | | |
| - | Box reference I.28 in case of Species: | select from the following: Aves, Ru Suidae, Pesca, Reptilian. | iminantia, Suidae, Mammalia | other than Ruminantia or | | | |
| Part II: | | | | | | | |
| (2) | Delete as a | ppropriate. | | | | | |
| - | The signat | ure and the stamp must be in a diffe | fferent colour to that of the printing. | | | | |
| - | this certifi | ne person responsible for the consig cate is only for veterinary purposes ntrol post of Great Britain, Channel I | and must accompany the con | | | | |
| Certifying O Name (in ca Date of signa Stamp | pital letters) | | Qualification and title Signature | | | | |
| | | | | | | | |
| | | | | | | | |
| | | | | | | | |
| | | | | | | | |
| | | | | | | | |