Veterinary certificate to EU

| 1.1. Consignor Name | | 1.2. Certificate reference number | | 1.2.a. | | |
|---|--|--|---------------|-----------------------------|------|--|
| Address | I.3. Central Competent Authority | | | | | |
| | DEPARTMENT FOR ENVIRONMENT, FOOD AND RURAL AFFAIRS | | | | | |
| | I.4. Local Competent Authority | | | | | |
| Tel. | | | | | | |
| I.5. Consignee | | I.6. Person responsible for the consignment | ent in the EU | IJ | | |
| Name | | Name | | | | |
| Address | | Address | | | | |
| | | | | | | |
| Postal Code | | Postal Code | | | | |
| Tel. I.7. Country of origin, ISO code | I & Design of origin Code | Tel I.9. Country of destination ISO code I.10. Region of destination Code | | | | |
| UNITED KINGDOM GB | I.8. Region of origin, Code | I.9. Country of destination | 180 code | 1.10. Region of destination | Code | |
| I.11. Place of origin | | I.12. Place of destination | | | | |
| Name | Approval number | | Custom v | warehouse | | |
| Address | | Name | | | | |
| | | Address | | | | |
| | | | | | | |
| | | | | | | |
| | | Postal Code | | | | |
| | | Approval number | | | | |
| I.13. Place of loading | | I.14. Date of departure | | | | |
| I.15 Means of transport | | VACE - DID : EV | | | | |
| | hin Pailway wasan | I.16. Entry BIP in EU | | | | |
| Road vehicle | hip Railway wagon Other | | | | | |
| Identification: | Other | I.17. | | | | |
| Documentary references: | | | | | | |
| I.18. Description of commodity | \wedge | I.19. Commodit | ty code (HS | code) | | |
| | | | | | | |
| | | | | | | |
| I.21. Temperature of products | 1.20 | Quantity | I.22. N | lumber of packages | | |
| Ambient Chilled | Frozen | | | | | |
| I.23. Seal/Container No. | | | I.24. T | ype of packaging | | |
| I . | | | | | | |
| =" | | | | | | |
| I.25. Commodity certified for: | | | | | | |
| 1.25. Commodity certified for: Technical use | | | | | | |
| | 1.27 | For import or admission into EU | | | | |
| Technical use | 1.27 | For import or admission into EU | | | | |
| Technical use | ISO code | For import or admission into EU | | | | |
| Technical use 1.26. For transit to 3rd country by EU | | For import or admission into EU | | | | |
| Technical use 1.26. For transit to 3rd country by EU | | For import or admission into EU | | | | |
| Technical use 1.26. For transit to 3rd country by EU 3rd country | ISO code | For import or admission into EU | | | | |
| Technical use 1.26. For transit to 3rd country by EU 3rd country | ISO code | | | Batch number | | |
| Technical use 1.26. For transit to 3rd country by EU 3rd country 1.28. Identification of the commodities | ISO code | number of establishments | | Batch number | | |
| Technical use 1.26. For transit to 3rd country by EU 3rd country 1.28. Identification of the commodities | ISO code | number of establishments | | Batch number | | |
| Technical use 1.26. For transit to 3rd country by EU 3rd country 1.28. Identification of the commodities | ISO code | number of establishments | | | | |
| Technical use 1.26. For transit to 3rd country by EU 3rd country 1.28. Identification of the commodities | ISO code | number of establishments | | | | |
| Technical use 1.26. For transit to 3rd country by EU 3rd country 1.28. Identification of the commodities | ISO code | number of establishments | | | | |
| Technical use 1.26. For transit to 3rd country by EU 3rd country 1.28. Identification of the commodities | ISO code | number of establishments | | | | |
| Technical use 1.26. For transit to 3rd country by EU 3rd country 1.28. Identification of the commodities | ISO code | number of establishments | | Batch number | | |
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| Technical use 1.26. For transit to 3rd country by EU 3rd country 1.28. Identification of the commodities | ISO code | number of establishments | | | | |
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| Technical use 1.26. For transit to 3rd country by EU 3rd country 1.28. Identification of the commodities | ISO code | number of establishments | | | | |
| Technical use 1.26. For transit to 3rd country by EU 3rd country 1.28. Identification of the commodities | ISO code | number of establishments | | | | |
| Technical use 1.26. For transit to 3rd country by EU 3rd country 1.28. Identification of the commodities | ISO code | number of establishments | | | | |
| Technical use 1.26. For transit to 3rd country by EU 3rd country 1.28. Identification of the commodities | ISO code | number of establishments | | | | |
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| Technical use 1.26. For transit to 3rd country by EU 3rd country 1.28. Identification of the commodities | ISO code | number of establishments | | | | |
| Technical use 1.26. For transit to 3rd country by EU 3rd country 1.28. Identification of the commodities | ISO code | number of establishments | | | | |
| Technical use 1.26. For transit to 3rd country by EU 3rd country 1.28. Identification of the commodities | ISO code | number of establishments | | | | |

en 1/4

Treated blood products, excluding those of equidae, for the manufacture of derived products for purposes outside the feed chain for farmed animals

| | II. Health information | | | II.a. Certificate reference number | II.b. | | | |
|------------------------|---------------------------|--|---|--|--|--|--|--|
| | TI. TICALLI III OTTIMATOR | | | | | | | |
| | | the European Pa | rliament and of the Council (1a), ar | t I have read and understood Regund in particular Article 8(c) and Artic (c)), and in particular Chapter II of A | le 8(d) and Article 10 thereof, | | | |
| tion | II.1. | the blood products described above consist of blood products that satisfy the requirements below; | | | | | | |
| fica | II.2. | | | | | | | |
| Part II: Certification | II.3. | | | | | | | |
| Part II | | | | | | | | |
| | | (²) and/or [- | with Union legislation, but which animals, derived from carcases | which is rejected as unfit for human did not show any signs of diseases is that have been slaughtered in mption following an ante-mortem in | communicable to humans or a slaughterhouse and were | | | |
| | | (²) and/or [- | humans or animals, obtained fro | which did not show any signs of m animals that have been slaughte human consumption following ar n;] | red in a slaughterhouse after | | | |
| | | (²) and/or [- | blood and blood products origin disease communicable through t | nating from live animals that did no hese products to humans or animal | ot show clinical signs of any s;] | | | |
| | | (²) and/or [- | blood and blood products de consumption;] | rived from the production of pro | oducts intended for human | | | |
| | | (²) and/or [- | animal by-products which have treatment as defined in Article 1 Directive 96/23/EC (2b);] | been derived from animals which h (2)(d) of Council Directive 96/22/EC | ave been submitted to illegal (^{2a}) or Article 2(b) of Council | | | |
| | | (²) and/or [- | listed in Group B(3) of Annex | residues of other substances and to Directive 96/23/EC, if such retion or, in the absence thereof, in na | sidues exceed the permitted | | | |
| | II.4. | accordance with | Union legislation, in slaughterhou tion or from live animals in faciliti | ed from was been collected in s ses approved and supervised by the es approved and supervised by the | ne competent authority of the | | | |
| | (²) [II.5. | In the case of blood products derived from Artiodactyla, Perissodactyla and Proboscide crossbreeds, other than Suidae and Tayassuidae, the products have undergone one of the folk guaranteeing the absence of pathogens of foot-and-mouth disease, vesicular stomatitis, rinderpes ruminants, Rift Valley fever and bluetongue: | | | | | | |
| | | (2) either [heat treatment at a temperature of 65 °C for at least three hours, followed by an effectiven check;] (2) and/or [irradiation at 25 kGy by gamma rays, followed by an effectiveness check;] | | | | | | |
| | | | | | | | | |
| | | (²) and/or | [change in pH to pH 5 for two ho | urs, followed by an effectiveness ch | neck;] | | | |
| | | (²) and/or | [heat treatment of at least 80 check.]] | °C throughout their substance, for | ollowed by an effectiveness | | | |

COUNTRY: UNITED KINGDOM

Treated blood products, excluding those of equidae, for the manufacture of derived products for purposes outside the feed chain for farmed animals

| II. Health information | | | | | II.a. Certificate reference i | number | II.b. | |
|------------------------|---|---|--|---|--|---|---|---|
| | | | | | | | | |
| | | | | | | | | |
| (²) [II.6. | (²) [II.6. In the case of blood products derived from Suidae, Tayassuidae, poultry and other avian species, the product undergone one of the following treatments guaranteeing the absence of pathogens of the following disease and-mouth disease, vesicular stomatitis, swine vesicular disease, classical swine fever, African swine Newcastle disease and highly pathogenic avian influenza, as appropriate to the species: | | | | | | eases: foot- | |
| | (²) either | [heat treatme check;] | ent at a tem | nperature o | of 65 °C for at lea | st three hours, f | ollowed by an ef | fectiveness |
| | (²) and/or | [irradiation at | 25 kGy by | gamma ra | ys, followed by ar | effectiveness ch | neck;] | |
| | (²) and/or | | [heat treatment of at least 80 °C for Suidae/Tayassuidae (2) and at least 70 °C for poultry and other avian species (2) throughout the substance of the product, followed by an effectiveness check]]. | | | | | |
| (²) [II.7. | | ood products derived from species other than those listed in point II.5 or II.6, the products have following treatment (please specify):] | | | | | | |
| II.8. | The products we | ere: | | | | | | |
| | (²) either [packed in new or sterilised bags or bottles,] | | | | | | | |
| | (²) or | | vere thoroughly o efore use;] and | leaned and | | | | |
| | the outer packag | ging or containers bear labels indicating 'NOT FOR HUMAN OR ANIMAL CONSUMPTION'; | | | | | | N'; |
| II.9. | the products we | re stored in encl | e stored in enclosed storage; | | | | | |
| II.10. | all precautions w | vere taken to avoid the contamination of the products with pathogenic agents after treatment; | | | | | | |
| (²) [II.11. | The treated bloc | od products described above | | | | | | |
| | (²) either | [is derived fro | om other rui | minants th | an bovine, ovine o | or caprine animal | s.]] | |
| | (²) or | [is derived fro | om bovine, | ovine or ca | prine animals and | d does not contai | n and is not deriv | ed from: |
| | | (²) either | continuo | usly reared | caprine materials d and slaughtered in accordance wit | in a country or r | egion classified | |
| | | (²) or | | | k material as defi 1 of the European | | | |
| | | | ca re n | aprine anii eared and egligible | y separated mea mals, except from slaughtered in a BSE risk in C (⁴), in which the | those animals to a country or reg accordance wi | hat were born, c gion classified a th Commission | ontinuously s posing a n Decision |
| | | | c th in in c | aprine aniline central strument into the continuously | product or derive mals which have nervous tissue ntroduced into the ranial cavity, ex y reared and slau gligible BSE risk in | been killed, afte by means of e cranial cavity, of cept for those aghtered in a cou | er stunning, by la an elongated or by means of g animals that n untry or region c | aceration of rod-shaped gas injected were born, classified as |

COUNTRY: UNITED KINGDOM

Treated blood products, excluding those of equidae, for the manufacture of derived products for purposes outside the feed chain for farmed animals

| | | the feed | a chain for farmed animals |
|-------------------|---|--|---|
| II. Health inform | mation | II.a. Certificate reference number | II.b. |
| | | | |
| Not | tes | | |
| Par | rt I: | | |
| _ | Box reference I.6: Person responsible for the consignment it is a certificate for a commodity to be transited through the commodity to be imported into the European Union. | | |
| _ | Box reference I.11 and I.12: Approval number: the regist issued by the competent authority. | tration number of the establishment | or plant, which has been |
| _ | Box reference I.12: Place of destination: this box is to be fi in transit may only be stored in free zones, free warehouse | | ansit commodity. Products |
| + | Box reference I.15: Registration number (railway wagons of is to be provided. In the case of unloading and reloading entry into the European Union. | | |
| _ ` | Box I.19: use the appropriate Harmonized System (HS) coo | de under the following headings: 05.1 | 1, 30.02, 35.02 or 35.04. |
| _ | Box reference I.23: for bulk containers, the container numb | er and the seal number (if applicable) | must be included. |
| _ | Box reference 1.25: technical use: any use other than production or manufacturing of pet food. | feeding of farmed animals, other th | nan fur animals, and the |
| _ | Box reference I.26 and I.27: fill in according to whether it is | a transit or an import certificate. | |
| _ | Box reference I.28 in case of Species, select from the Ruminantia or Suidae, Pesca, Reptilian. | following: Aves, Ruminantia, Suida | e, Mammalia other than |
| Par | rt II: | | |
| (^{1a}) | OJ L 300, 14.11.2009, p. 1. | | |
| (^{1b}) | OJ L 54, 26.2.2011, p. 1. | | |
| (²) | Delete as appropriate. | | |
| (^{2a}) | OJ L 125, 23.5.1996, p. 3. | | |
| (^{2b}) | OJ L 125, 23.5.1996, p. 10. | | |
| (³) | OJ L 147, 31.5.2001, p. 1. | | |
| (4) | OJ L 172, 30.6.2007, p. 84. | | |
| _ | The signature and the stamp must be in a different colour to | o that of the printing. | |
| _ | Note for the person responsible for the consignment in the and must accompany the consignment until it reaches the based on the second of the | European Union: this certificate is or porder inspection post of the European | lly for veterinary purposes n Union. |
| | | | |
| | | | |
| | | | |
| | | | |
| Official ve | eterinarian/Official inspector | | |
| Nam | ne (in capital letters): | Qualification and title: | |
| Date | e: | Signature: | |
| Stan | mp: | | |
| | | | |

en 4/4