| | I.1. Consignor | | | | I.2. IMSOC Ref | erence | | | |
|---------------------------------|-----------------------|-------------------------|---|-------------------------------------|---------------------------|---------------------------------|----------|--------------------|--------------|
| | Name | | | | I.2.a. Local Reference | | | | |
| | Address | | | | | | | | |
| | Country ISO Code | | | | | | | | |
| | I.5. Consignee | | | | I.3. Central co | mpetent authority | | | |
| 님 | Name | | | | I.4. Local com | petent authority | | | |
| ler | Address | | | | | | | | |
| E | Country | | ISO Code | | | | | | |
| Part I : Details of consignment | I.7. Country of orig | rin | | ISO Code | I.9. Country of | destination | | | ISO Code |
| ü | | · | | | | | | | |
| Ĺ C | I.8. Region of origi | n | | Code | I.10. Region of | destination | | | |
| S O | I.11. Place of Dispa | | | coue | I.12. Place of d | | | | |
| tail | Name | | | | Name | | | | |
| Qet | Address | | | | Address | | | | |
| $\overline{}$ | Approval Number | 2 | | | Approval Nui | nber | | | |
| ะเ | Country | | ISO Code | | Country | | | ISO Code | |
| Pa | I.13. Place of Loadi | ing | | | I 14 Data and | time of departure | | | |
| | | шід | | | 1.14. Date anu | unie of departure | | | |
| | Name Address | | | | | | | | |
| | Approval Number | | | | | | | | |
| | Country | - | ISO Code | | | | | | |
| | - | | | | | | | | |
| | I.15. Means of Tran | - | | | I.16 Entry Poin | nt | | | |
| | Mode | International transport | Identification | | | | | | |
| | | document | | | | | | | |
| | | | | | | | | | |
| | | | | | | | | | |
| | | | | | | | | | |
| | I.18. Transport cor | ditions | | | I 17 Accompa | nying documents | | | |
| | Frozen | Chilled 🗆 | Ambient 🛛 🛛 Co | ntrolled | Accompanyi | itying documents | | | |
| | temperature | | | ng | ng Date of issue | | | | |
| | | | | document Duce of Issue reference | | | | | |
| | | | | | Country | | Place of | of | |
| | | | | | | issue | | | |
| | I.19. Container No | / Seal No | | | | | | | |
| | I.20. Certified as | | | | | | | | |
| | Human consumpti | | Technical use 🛛 | | Other Animal Feedingstuff | | | | |
| | Breeding and prod | | Production 🗆 | | | Production of petfood Slaughter | | _ | |
| | Pharmaceutical us | e 🗀 | Breeding \Box | | Relaying 🗆 | | Artific | ial reproductio | n 🗀 |
| | Fattening 🗆 | | | | | | | | |
| | I.21. For transit thi | rough a third cou | ntry 🗌 | | I.22. For trans | it through Member Sta | te(s) | | |
| | Country | | ISO Code | | | 0 | | | |
| | EU Exit | | BCP code | | | | | | |
| | Authority | | | | Country | | ISO Co | de | |
| | EU Entry Authority | | BCP code | | | | | | |
| | I.23. Total number | of packages | I.24. Total quantity | | I.25. Total net | weight | I.25. T | otal gross weigl | nt |
| | I.28. Description of | fconsignment | | | | | | | |
| | - | - | FLOUR, STARCH OR MI | I V. DASTRVCO | | | | | |
| | | | | | | | taining | less than 401% | hy weight of |
| | cocoa calculate | d on a totally defa | ions of flour, groats, me atted basis, not elsewhe ess than 5 % by weight | re specified or i | ncluded; food p | reparations of goods of con | f headi | ngs 0401 to 04 | 04, not |
| | | | | | aleu oli a totaliy | | ewitere | | |
| | Commodity | Speci | es | Quantity | | Net weight | | Package count | |
| | Identification number | | | | | | | | |
| | | | | Identification | system | | | | |
| | | | | | | | | | |
| | | | | | | | | | |
| | | | | | | | | | |
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| | | | | | | | | | |

| | II. Health information | | | | | | |
|---|--|--|---|---|---|------------------------------------|---|
| | I, the undersigned offi above contain: | cial veterina | rian/officia | l inspector h | ereby certify that the | e compos | ite products described |
| | (1) either □ II.1.A Meat products, treated stomachs, bladders and intestines (2) in any quantity and such meat products, treated stomachs, bladders and intestines have been produced according to Commission Decision 2007/777/EC and contain the following meat constituents and meet the criteria indicated below | | | | | | |
| cation | Species (A) | Treatment (B) | Origin (C) | | | | |
| Part II: Certification | (A) | (A) Insert the code for the relevant species of meat product, treated stomachs, bladders and intestines where BOV = domestic bovine animals (Bos Taurus, Bison bison, Bubalus bubali and their cross breeds); OVI = domestic sheep (Ovis aries) and goats (Capra hircus); EQI = domestic equine animals (Equus caballus, Equus asinus and their cross breeds), POR=domestic porcine animals (Sus scrofa); RM = Domestic rabbits, PFG = domestic poultr and farmed feather game, RUF = farmed non-domestic animals other than suidae and solipeds; RUW = wild non-domestic animals other than suidae and solipeds; SUW = wild non-domestic solipeds, WL = wild lagomorphs, WGB = wild game bids | | | | | ison bison, Bubalus bubalis oats (Capra hircus); EQI = ir cross breeds), oits, PFG = domestic poultry other than suidae and nd solipeds; SUW = wild non- |
| | (B) | | | for the requ 2007/777/EC | | ecified ar | nd defined in Parts 2, 3 and 4 |
| | (C) | and intesti regionaliza indicated i | nes as listed ition by reta n Part 1 of A | l in Annex 2, ained EU law Annex 2 to D | Part 2 to Decision 20 for the relevant mea | 07/777/E at constit | |
| | | | - | | s the country of expo | rt in box | I.7, |
| | | | - | Great Brita | | | , |
| - a third country or parts thereof authorised to export to meat products treated with treatment A as set out in Ar Decision 2007/777/EC, where the third country where th product is produced is also authorised to export to Grea meat products treated with that treatment. | | | | | as set out in Annex 2 to ountry where the composite o export to Great Britain | | |
| (1) and/or \Box [II.1.B Processed dairy products (3) in an amount of half or more of the substance of the c product or not shelf stable dairy products in any quantity that | | | | | bstance of the composite | | |
| | | (a) | | - | the country one of the following | | ne country of origin of the |
| | | | - | the same as | s the country of expo | rt in box | I.7, |
| | | | - | Great Brita | | | |
| | | | - | products in 605/2010, w produced is | Column A or B of An where the third count | inex 1 to ry where er the sa | reat Britain milk and dairy Regulation (EU) No e the composite product is me conditions, to export to |
| The country of origin indicated in box I.7 must be listed in Annex 1 to Regulation (EU) 605/2010 and the treatment applied must conform to the treatment provided for in th for the relevant country; | | | | | - | | |
| | | (b) | have been | produced fr | om milk obtained fro | m anima | als: |
| | | | (i) | under the o | ontrol of the official | veterina | ry service; |
| | | | (ii) | | o holdings which we disease or rinderpes | | nder restrictions due to foot- |
| | | | (iii) | the animal | health conditions lai | d down i | to ensure that they satisfy n Chapter I of Section IX of and in Directive 2002/99/EC |
| | | (c) | are dairy p | products mad | le from raw milk obt | ained fro | om |

| E | UROPEAN | UNION | (GB) Composite j | products transit or stora | ge from EU 28/2012 GBHC089E | |
|---|---|---|---------------------------|---|--|--|
| | II. Health in | formation | | | | |
| | (1) | | Great Bri | | loes and prior to import into en produced from raw milk | |
| Part II: Certification | (1) | | | heat treatment with a he that achieved by a paste for 15 seconds and when | on treatment involving a single eating effect at least equivalent to urisation process of at least 72°C re applicable, sufficient to ensure n alkaline phosphatase test er the heat treatment;] | |
| II: Cei | (1) | | | or \circ [a sterilisation proctor to or greater than three; | ess, to achieve an F0 value equal] | |
| Part | (1) | | | | perature (UHT) treatment at not ination with a suitable holding | |
| | (1) or \circ [a high temperature short time pasteurist treatment (HTST) at 72°C for 15 seconds, or a with an equivalent pasteurisation effect, appl with a pH lower than 7,0 achieving, where ap negative reaction to an alkaline phosphatase | | | | | |
| | (1) or ○ [a high temperature short time pasteurisation treatment at 72°C for 15 seconds, or a treatment with an equivalent pasteurisation effect, applied twice to milk with a pH equal to great than 7,0 achieving where applicable, a negative reaction alkaline phosphatase test, immediately followed by | | | | | |
| | (1) | | | either \circ [lowering the p | H below 6 for one hour;] | |
| | (1) | | | or \circ [additional heating combined with desiccation ${f c}$ | g equal to or greater than 72°C, ion;]] | |
| | (1) | | import ir | | s, goats or buffaloes and prior to ergone or been produced from | |
| | (1) | | | either \circ [a sterilisation equal to or greater than | process, to achieve an F0 value three;] | |
| | | | | perature (UHT) treatment at not ination with a suitable holding | | |
| | | (d) | were produced on (4).] | or between | and | |
| | (5) | | | | | |
| Were produced from eggs coming from an establishment which satisfies the requirement of Annex 3 to Regulation (EC) No 853/2004 which at the date of issue of the certificate is fr pathogenic avian influenza as defined in Regulation (EC) No 798/2008 and | | | | | | |
| | either | | | | | |
| | (1) □ [II.1.C.1 [within a 10km radius of which [including, where appropriate, the territory of a neighbouring country,] there has been no outbreak of highly pathogenic avian influenza or Newcas disease for at least the previous 30 days.] or (1) □ [II.1.C.2 [the egg products were processed: | | | | | |
| | | | | | | |
| | (1) | either \circ [liquid egg white was treated: | | | | |

- either \circ [liquid egg white was treated: (1)
- (1) either o [with 55.6°C for 870 seconds.]
- (1) or \circ [with 56.7°C for 232 seconds.]
- or $~\circ$ [10% salted yolk was treated with 62.2°C for 138 seconds.] (1) (1)
 - or \circ [dried egg white was treated:
- (1) either o [with 67°C for 20 hours]

(GB) Composite products transit or storage from EU 28/2012 GBHC089E

| | II. Health information | | | |
|---------------|------------------------|---|---------------|--|
| | (1) | or o [with 54.4°C for 53 | 12 hours ll | |
| | (1) | $01 \circ [\text{with } 34.4 \circ 101 \circ]$ | 15 110015.]] | |
| | (1) | or \circ [whole eggs were at least trea | ted: | |
| | (1) | either \circ [with 60°C for | 188 seconds.] | |
| | (1) | or \circ [completely cooke | ed.] | |
| ទ | | \circ [whole egg blends were at least | treated]: | |
| Certification | (1) | either \circ [with 60°C for | 188 seconds.] | |
| tifi | (1) | or \circ [with 61.1°C or 94 | seconds.] | |
| G | | | | |
| H | | | | |
| Part | | | | |

| | II. Health info | rmation | | | | | | | |
|--|--|---|---|--|---|--|--|--|--|
| | | | | | | | | | |
| | Notes | Jotes | | | | | | | |
| | | *) Those countries subject to the transitional import arrangements include: an EU member State; Iceland; | | | | | | | |
| | | | and Switzerland. | | | | | | |
| cation | References to Great Britain in this certificate include Channel Islands and Isle of Man. 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). | | | | | | | | |
| Ē | Part I: | | - | | | | | | |
| Part II: Certification | — | Box reference I.7: | Insert ISO code of the country of origin of the meat product, treated stomachs, bladders and intestines as listed in Annex 2, Part 2 to Decision 2007/777/EC and/or for processed dairy product in Annex 1 to Commission Regulation (EU) No 605/2010. | | | | | | |
| Ğ | _ | Box Reference I.11: | ference country of origin which must be the same as the country of origin inbox I.7. | | | | | | |
| | | Approval number is not applicable. | | | | | | | |
| | | Box reference I.15: | (aircraft) or name (ship). In the cas containers and their registration n | ons or container and road vehicles), flight number se of transport in containers, the total numbers of number and where there is a serial number of the seal it ase of unloading and reloading, the consignor must inform tion into Great Britain. | | | | | |
| | _ | Box I.16: | Do not use this box until the end of | f the transitional staging perio | od. | | | | |
| | _ | Box reference I.19: | For containers or boxes, the container number and the seal number (if applicable) must be | | | | | | |
| | _ | Box Reference I.25: | Indicate total gross weight and tota | ıl net weight. | | | | | |
| | Box Use the appropriate Harmonised S reference as: 16.01; 16.02; 16.03; 16.04; 16.05; I.28: | | | | | | | | |
| | _ | Box reference I.28: | Manufacturing Plant: insert the na establishments of production of the composite products containing me indicate "meat product", "treated s product containing dairy products | e composite product(s). Natur at products, treated stomachs tomachs", "bladders" or "inte | e of commodity: in case of , bladders and intestines | | | | |
| | Part II: | | | | | | | | |
| | (1) | Keep as ap | propriate. | | | | | | |
| | (2) | 2) Meat products as laid down in point 7.1 of Annex 1 to Regulation (EC) No 853/2004 and treated stomachs, bladders and intestines as laid down in point 7.9 of Annex 1 to Regulation (EC) No 853/2004 that have undergone one of the treatments laid down in Annex 2 part 4 to Decision 2007/777/EC. | | | | | | | |
| Raw milk and dairy products means, raw milk and dairy products for human consumption point 7.2 of Annex 1 to Regulation (EC) No 853/2004. | | | | | n consumption as defined in | | | | |
| | (4) | 4) Date or dates of production. Imports of raw milk and dairy products shall not be allowed when obtained either prior to the date of authorisation for exportation to Great Britain of the third country or part thereof mentioned under I.7 and I.8 or during a period where restrictive measures have been adopted by Great Britain against imports of raw milk and dairy products from this third country or part thereof. | | | | | | | |
| | (5) | Country of origin authorised to export to Great Britain. | | | | | | | |
| The colour of the signature shall be different to that of the printing. The same rule applies to state other than those embossed or watermark. | | | | | ne rule applies to stamps | | | | |
| | Certifying Officer | | | | | | | | |

(GB) Composite products transit or storage from EU 28/2012 GBHC089E

| II. Health information | | |
|---|--------------------------------------|--|
| Name (in capital letters) Date of signature Stamp | Qualification and title Signature | |
| | | |