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►<u>C1</u> COMMISSION REGULATION (EU) No 206/2010

of 12 March 2010

laying down lists of third countries, territories or parts thereof authorised for the introduction into the European Union of certain animals and fresh meat and the veterinary certification requirements

(Text with EEA relevance) ◀

(OJ L 73, 20.3.2010, p. 1)

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► <u>M3</u>	Commission Implementing Regulation (EU) No 342/2011 of 8 April 2011	L 96	10	9.4.2011
► <u>M4</u>	Commission Implementing Regulation (EU) No 801/2011 of 9 August 2011	L 205	27	10.8.2011

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- ►<u>C1</u> Corrigendum, OJ L 146, 11.6.2010, p. 1 (206/2010)
- ►<u>C2</u> Corrigendum, OJ L 49, 24.2.2011, p. 53 (144/2011)
- ►<u>C3</u> Corrigendum, OJ L 63, 10.3.2011, p. 28 (144/2011)

COMMISSION REGULATION (EU) No 206/2010

of 12 March 2010

laying down lists of third countries, territories or parts thereof authorised for the introduction into the European Union of certain animals and fresh meat and the veterinary certification requirements

(Text with EEA relevance)

THE EUROPEAN COMMISSION,

Having regard to the Treaty on the Functioning of the European Union,

Having regard to Council Directive 92/65/EEC of 13 July 1992 laying down animal health requirements governing trade in and imports into the Community of animals, semen, ova and embryos not subject to animal health requirements laid down in specific Community rules referred to in Annex A (I) to Directive 90/425/EEC (1), and in particular Articles 17(2)(b) and 17(3)(a), the first subparagraph of Article 17(3)(c), the fourth indent of Article 18(1) and Article 19 thereof,

Having regard to Council Directive 2002/99/EC of 16 December 2002 laying down the animal health rules governing the production, processing, distribution and introduction of products of animal origin for human consumption (2), and in particular Article 8, Article 9(2)(b) and Article 9(4) thereof,

Having regard to Council Directive 2004/68/EC of 26 April 2004 laying down animal health rules for the importation into and transit through the Community of certain live ungulate animals, amending Directives 90/426/EEC and 92/65/EEC and repealing Directive 72/462/EEC (3), and in particular the first and second subparagraphs of Article 3(1), the first subparagraph of Article 6(1), Article 7(e), Article 8, the first paragraph of Article 10 and Article 13(1) thereof,

Having regard to Regulation (EC) No 852/2004 of the European Parliament and of the Council of 29 April 2004 on the hygiene of foodstuffs (4), and in particular Article 12 thereof,

Having regard to Regulation (EC) No 853/2004 of the European Parliament and of the Council of 29 April 2004 laying down specific hygiene rules for food of animal origin (5), and in particular Article 9 thereof,

⁽¹⁾ OJ L 268, 14.9.1992, p. 54.

⁽²⁾ OJ L 18, 23.1.2003, p. 11. (3) OJ L 139, 30.4.2004, p. 321.

⁽⁴⁾ OJ L 139, 30.4.2004, p. 1.

⁽⁵⁾ OJ L 139, 30.4.2004, p. 55.

Having regard to Regulation (EC) No 854/2004 of the European Parliament and of the Council of 29 April 2004 laying down specific rules for the organisation of official controls on products of animal origin intended for human consumption (1), and in particular Article 11(1) and Article 16 thereof,

Having regard to Regulation (EC) No 882/2004 of the European Parliament and of the Council of 29 April 2004 on official controls performed to ensure the verification of compliance with feed and food law, animal health and animal welfare rules (2), and in particular Article 48(1) thereof,

Whereas:

- (1) Council Directive 72/462/EEC of 12 December 1972 on health and veterinary inspection problems upon importation of bovine, ovine and caprine animals and swine, fresh meat or meat products from third countries (3) provided for a list to be drawn up of the countries or parts thereof from which Member States are to authorise the importation of certain live animals and fresh meat of certain animals.
- (2) Accordingly, Council Decision 79/542/EEC of 21 December 1976 drawing up a list of third countries or parts of third countries, and laying down animal and public health and veterinary certification conditions, for importation into the Community of certain live animals and their fresh meat (4) was adopted. That Decision establishes the sanitary conditions for the importation into the European Union of live animals excluding equidae, and for the importation of fresh meat of such animals, including equidae, but excluding meat preparations. Annexes I and II to that Decision also set out lists of third countries or parts thereof from which certain live animals and their fresh meat may be imported into the Union as well as models of veterinary certificates.
- (3) Since the date of adoption of that Decision, a number of new animal health and public health requirements have been laid down in other Union acts, constituting a new regulatory framework in this area. Also, Directive 72/462/EEC has been repealed by Directive 2004/68/EC.
- (4) Article 20 of Directive 2004/68/EC states that implementing rules on import established in accordance with Decisions adopted pursuant to Directive 72/462/EEC, inter alia Decision 79/542/EEC, shall remain in force until replaced by measures adopted under the new regulatory framework.

⁽¹⁾ OJ L 139, 30.4.2004, p. 206.

⁽²⁾ OJ L 165, 30.4.2004, p. 1.

⁽³⁾ OJ L 302, 31.12.1972, p. 28.

⁽⁴⁾ OJ L 146, 14.6.1979, p. 15.

- (5) In accordance with Article 4(3) of Directive 2004/41/EC of the European Parliament and of the Council of 21 April 2004 repealing certain Directives concerning food hygiene and health conditions for the production and placing on the market of certain products of animal origin intended for human consumption and amending Council Directives 89/662/EEC and 92/118/EEC and Council Decision 95/408/EC (¹), once the necessary provisions on the basis of Regulations (EC) No 852/2004, (EC) No 853/2004, (EC) No 854/2004 or Directive 2002/99/EC are adopted, the implementing rules adopted on the basis of Directive 72/462/EEC shall cease to apply.
- (6) Decision 79/542/EEC has been amended several times and import provisions based on the new regulatory framework have already been introduced in Decision 79/542/EEC. For the sake of clarity and transparency the measures that are laid down in Decision 79/542/EEC should be laid down in a new legal act. This Regulation includes all the provisions of Decision 79/542/EEC. Consequently, by the entry into force of the present Regulation Decision 79/542/EEC is lapsed and thus no longer applies, pending the explicit repeal of it.
- (7) Directive 92/65/EEC lays down the animal health requirements governing trade in and imports into the Union of live animals, semen, ova and embryos not subject to the animal health requirements laid down in the specific Union acts referred to in Annex F to that Directive. Pursuant to that Directive, those live animals, semen, ova and embryos may be imported into the Union only from a third country which is on a list drawn up in accordance with the procedure referred to in that Directive. In addition, such live animals are to be accompanied by a health certificate corresponding to a specimen drawn up in accordance with the procedure referred to therein.
- (8) Council Directive 96/93/EC of 17 December 1996 on the certification of animals and animal products (²) lays down the rules to be observed in issuing the certificates required by veterinary legislation to prevent misleading or fraudulent certification. It is appropriate to ensure that rules and principles at least equivalent to those laid down in that Directive are applied by the official inspectors or veterinarians of third countries. Certain third countries, which are listed in Annex II to this Regulation, have provided sufficient guarantees as to the existence and implementation of such rules and principles. It is therefore appropriate to authorise the introduction of certain live animals into the Union from those third countries, provided that no further restrictions are required by their specific disease situation.
- (9) Directive 2002/99/EC lays down the animal health rules concerning the introduction into the Union of products of animal origin and products obtained therefrom intended for human consumption. Pursuant to that Directive, lists are to be drawn up of the third countries or regions of third countries from which imports of specified products of animal origin are permitted and those imports are to comply with certain veterinary certification requirements.

⁽¹⁾ OJ L 157, 30.4.2004, p. 33.

⁽²⁾ OJ L 13, 16.1.1997, p. 28.

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- (10) Directive 2004/68/EC lays down the animal health requirements for the importation into and transit through the Union of certain live ungulates. The importation of those live ungulates into and their transit through the Union is authorised only from third countries and territories that appear on a list or lists drawn up in accordance with the procedure referred to in that Directive and those imports are to comply with certain veterinary certification requirements.
- (11) Save the provisions of article 17(2) last subparagraph of Directive 92/65/EEC, live animals, and products of animal origin to which Directives 92/65/EEC, 2002/99/EC and 2004/68/EC apply are to be imported into or transit through the Union only if they are accompanied by a veterinary certificate and comply with the relevant requirements laid down in Union legislation.
- (12) Accordingly, for the implementation of Directives 92/65/EEC, 2002/99/EC and 2004/68/EC, it is appropriate to lay down in this Regulation lists of third countries, territories and parts thereof and the specific import conditions including model veterinary certificates for certain live animals and the fresh meat of certain animals.
- In the interest of consistency of Union legislation, this Regulation should also take into account the public heath requirements laid down in other Union acts and in particular in Regulations (EC) Nos 852/2004, 853/2004 and 854/2004 which lay down rules concerning the hygiene of foodstuffs and food of animal origin and rules for the organisation of official controls on products of animal origin intended for human consumption, as well as the requirements of Council Directive 96/23/EC of 29 April 1996 on measures to monitor certain substances and residues thereof in live animals and animal products (1), and of Regulation (EC) No 999/2001 of the European Parliament and of the Council of 22 May 2001 laying down rules for the prevention, control and eradication of certain transmissible spongiform encephalopathies (2).
- (14) Regulation (EC) No 882/2004 lays down general rules governing the performance of official controls carried out in the areas of food and feed, animal health and animal welfare. Article 48 thereof empowers the Commission to adopt a list of third countries from which specific products may be imported into the Union. Regulation (EC) No 854/2004 provides specific rules for the organisation of official controls on products of animal origin intended for human consumption, including the establishment of lists of third countries from which imports of products of animal origin are permitted. Those rules provide that those lists may be combined with other lists drawn up for public and animal health purposes.

⁽¹⁾ OJ L 125, 23.5.1996, p. 10.

⁽²⁾ OJ L 147, 31.5.2001, p. 1.

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- (15) The model certificates set out in the Annexes to this Regulation should therefore include attestations certifying that the public health requirements laid down in Directive 96/23/EC and Regulations (EC) No 999/2001, 852/2004, 853/2004 and 854/2004, are fulfilled.
- (16) The model certificates set out in the Annexes to this Regulation should also include attestations certifying that animal welfare requirements laid down in Council Directive 93/119/EC of 22 December 1993 on the protection of animals at the time of slaughter and killing (¹) and Council Regulation (EC) No 1/2005 of 22 December 2004 on the protection of animals during transport and related operations (²) are fulfilled.
- (17) In order to ensure that the health of live animals introduced into the Union is not jeopardised during their transport from the third country of origin to the Union, certain requirements relating to the transport of live animals should be laid down, including requirements on assembly centres.
- (18) In the interest of ensuring the protection of animal health in the Union, live animals should be conveyed directly to their place of destination in the Union.
- (19) Fresh meat introduced into the Union for transit to another third country poses a negligible risk to public health. Such meat should, however, comply with all the relevant animal health requirements. Accordingly, specific provisions on the transit, and storage before transit, of fresh meat should therefore be laid down.
- (20) Specific conditions for transit via the Union of consignments to and from Russia should be provided for owing to the geographical situation of Kaliningrad which affects only Latvia, Lithuania and Poland.
- (21) Consignments of fresh meat, excluding offal and minced meat, of farmed non-domesticated animals of the order Artiodactyla, originating from animals caught in the wild should be authorised for introduction into the Union. In order to rule out any possible animal health risks which could be posed by such introduction, it is appropriate that those animals be separated from wild animals for a period of three months prior to the introduction into the Union of such consignments. Accordingly, the model veterinary certificate for those consignments (RUF) should take that into account.
- (22) Commission Decision 2003/881/EC of 11 December 2003 concerning the animal health and certification conditions for imports of bees (*Apis mellifera* and *Bombus* spp.) from certain third countries (³) lays down the animal health and certification conditions for imports of bees from certain third countries. In the interest of simplification of Union legislation, the measures laid down in that Decision should be included in this Regulation. Consequently, Decision 2003/881/EC should be repealed.

⁽¹⁾ OJ L 340, 31.12.1993, p. 21.

⁽²⁾ OJ L 3, 5.1.2005, p. 1.

⁽³⁾ OJ L 328, 17.12.2003, p. 26.

- (23) It's appropriate to introduce a transitional period to allow Member States and industry to take the necessary measures to comply with the requirements laid down in this Regulation.
- (24) The measures provided for in this Regulation are in accordance with the opinion of the Standing Committee on the Food Chain and Animal Health,

HAS ADOPTED THIS REGULATION:

CHAPTER I

SUBJECT MATTER, SCOPE AND DEFINITIONS

Article 1

Subject matter and scope

- 1. This Regulation lays down the veterinary certification requirements for the introduction into the Union of consignments containing the following live animals or fresh meat:
- (a) ungulates;
- (b) the animals listed in Part 2 of Annex IV;
- (c) fresh meat intended for human consumption, excluding meat preparations, of ungulates and equidae.
- 2. This Regulation lays down the lists of third countries, territories or parts thereof from which the consignments referred to in paragraph 1 may be introduced into the Union.
- 3. This Regulation shall not apply to the introduction into the Union of non-domesticated animals:
- (a) for shows or exhibitions where such live animals are not regularly kept or bred;
- (b) forming part of circuses;
- (c) intended for an approved body, institute or centre as defined in Article 2(1)(c) of Directive 92/65/EEC.
- 4. This Regulation shall apply without prejudice to any specific certification requirements laid down in other Union acts or in agreements concluded by the Union with third countries.

Article 2

Definitions

For the purposes of this Regulation, the following definitions shall apply:

(a) 'ungulates' means ungulates as defined in Article 2(d) of Directive 2004/68/EC;

- (b) 'fresh meat' means fresh meat as defined in point 1.10 of Annex I to Regulation (EC) No 853/2004;
- (c) 'equidae' means equidae as defined in Article 2(b) of Council Directive 90/426/EEC (¹);
- (d) 'holding' means a farm or other officially supervised agricultural, industrial or commercial undertaking, including zoos, amusement parks and wildlife or hunting reserves where live animals are regularly kept or bred.

CHAPTER II

CONDITIONS FOR THE INTRODUCTION OF LIVE ANIMALS INTO THE UNION

Article 3

General conditions for the introduction of ungulates into the Union

Consignments of ungulates shall only be introduced into the Union if they comply with the following conditions:

- (a) they come from the third countries, territories or parts thereof listed in columns 1, 2 and 3 of the table set out in Part 1 of Annex I for which there is a model veterinary certificate corresponding to the consignment concerned listed in column 4 of the table in Part 1 of Annex I;
- (b) they are accompanied by the appropriate veterinary certificate, drawn up in accordance with the relevant model veterinary certificate set out in Part 2 of Annex I, taking into account the specific conditions indicated in column 6 of the table in Part 1 of that Annex, and completed and signed by an official veterinarian of the exporting third country;
- (c) they comply with the requirements set out in the veterinary certificate referred to in point (b), including:
 - (i) the supplementary guarantees laid down in that certificate, where indicated in column 5 of the table in Part 1 of Annex I;
 - (ii) any additional veterinary certification requirements that the Member State of destination may impose in accordance with Union veterinary legislation and which are included in the certificate.

Article 4

Conditions for assembly centres for certain consignments of ungulates

Consignments of ungulates which contain live animals from more than one holding shall only be introduced into the Union if they are assembled in assembly centres approved by the competent authority of the third country of origin in accordance with the requirements set out in Part 5 of Annex I.

Article 5

Protocols for the standardisation of materials and sampling and testing procedures for ungulates

Where sampling and testing is required by the veterinary certificates listed in column 4 of the table in Part 1 of Annex I for the diseases listed in Part 6 of that Annex, for the introduction into the Union of consignments of ungulates, such sampling and testing shall be carried out by or under the control of the competent authority of the third country of origin in accordance with the Protocols for the standardisation of materials and testing procedures set out in Part 6 of that Annex.

Article 6

Special conditions for certain consignments of ungulates imported into St Pierre and Miquelon and introduced into the Union

Consignments of ungulates of the species listed in the table in Part 7 of Annex I which were introduced into St Pierre and Miquelon less than six months prior to the date of shipment from St Pierre and Miquelon to the Union shall only be introduced into the Union if:

- (a) they comply with the residence and quarantine requirements set out in Chapter 1 of that Part;
- (b) they have been tested in accordance with the animal health test requirements set out in Chapter 2 of that Part.

Article 7

General conditions for the introduction into the Union of certain species of bees

- 1. Consignments of bees of the species listed in table 1 of Part 2 of Annex IV shall only be introduced into the Union from third countries or territories:
- (a) listed in Part 1 of Annex II;
- (b) where the presence of the American foulbrood, the small hive beetle (Aethina tumida) and the Tropilaelaps mite (Tropilaelaps spp.) is subject to compulsory notification throughout the whole territory of the third country or territory concerned.
- 2. By way of derogation from paragraph 1(a), consignments of bees may be introduced into the Union from a part of a third country or territory listed in Part 1 of Annex II which is:
- (a) a geographically and epidemiologically isolated part of the third country or territory
- (b) listed in the third column of the table in Section 1 of Part 1 of Annex IV.

When that derogation is applied, the introduction into the Union of consignments of bees shall be prohibited from all other parts of the third country or territory concerned not listed in the third column of the table in Section 1 of Part 1 of Annex IV.

- 3. Consignments of bees of the species listed in table 1 of Part 2 of Annex IV shall consist of either:
- (a) cages of queen bees (*Apis mellifera* and *Bombus* spp.), each containing one single queen bee with a maximum of 20 accompanying attendants; or
- (b) containers of bumble bees (*Bombus* spp.), each containing a colony of a maximum of 200 adult bumble bees.
- 4. Consignments of bees of the species listed in table 1 of Part 2 of Annex IV shall:
- (a) be accompanied by the appropriate veterinary certificate, drawn up in accordance with the relevant model veterinary certificate set out in Part 2 of Annex IV, and completed and signed by an official inspector of the exporting third country;
- (b) comply with the veterinary requirements set out in the veterinary certificate referred to in point (a).

Article 8

General conditions concerning the transport of live animals to the Union

During the period after loading in the third country of origin and before arrival at the border inspection post of introduction into the Union, consignments of live animals shall not be:

- (a) transported together with live animals that:
 - (i) are not intended for introduction into the Union; or
 - (ii) are of a lower health status;
- (b) unloaded in, or when transported by air, moved to another aircraft, or transported by road, by rail or moved on foot through a third country, territory or a part thereof which is not listed in columns 1, 2 and 3 of the table set out in Part 1 of Annex I or for which there is no model veterinary certificate corresponding to the consignment concerned listed in column 4 of the table in Part 1 of Annex I.

Article 9

Time limit for the period of transport to the Union of live animals

Consignments of live animals shall only be introduced into the Union where the consignment arrives at the border inspection post of introduction into the Union within 10 days of the date of issue of the appropriate veterinary certificate.

In the case of transport by sea, that period of 10 days shall be extended by an additional period corresponding to the duration of the journey by sea, as certified by a signed declaration of the master of the ship, drawn up in accordance with Part 3 of Annex I and attached in its original form to the veterinary certificate.

Article 10

Special conditions regarding the spraying of consignments of live animals transported by air to the Union

Where consignments of live animals, excluding consignments of bees, are transported by air, the crate or container in which they are transported and the surrounding area shall be sprayed with an appropriate insecticide.

That spraying shall be carried out immediately prior to the closing of the aircraft doors after loading, and after any subsequent opening of the doors in a third country, until the aircraft reaches its final destination.

The captain of the aircraft shall certify that the spraying has been carried out by signing a declaration, drawn up in accordance with Part 4 of Annex I and attached in its original form to the veterinary certificate.

Article 11

Conditions to be applied following the introduction into the Union of certain consignments of ungulates

1. Following their introduction into the Union, consignments of ungulates intended for breeding and production, or intended for zoos, amusement parks and wildlife or hunting reserves, shall be conveyed without delay to the holding of destination.

The ungulates shall remain on that holding for a period of at least 30 days, unless they are dispatched directly to a slaughterhouse.

2. Following their introduction into the Union, consignments of ungulates intended for immediate slaughter shall be conveyed without delay to the slaughterhouse of destination where they shall be slaughtered within five working days from the date of arrival at the slaughterhouse.

Article 12

Specific conditions concerning the transit through third countries of certain consignments of ungulates

Where specific condition I of Part 1 of Annex I applies, in order to allow consignments of the ungulates referred to in that condition originating in one Member State and destined for another Member State, to transit through a third country, territory or part thereof which is listed in the table in Part 1 of Annex I but for which there is no corresponding model veterinary certificate for consignments of the ungulates concerned indicated in column 4 of that table, the following conditions shall apply:

- (a) for bovine animals for fattening:
 - (i) the holdings of final destination must be designated in advance by the competent authority of the final destination;

- (ii) the live animals comprised in the consignment must not be moved from the holding of final destination unless for immediate slaughter;
- (iii) all movements of live animals into and out of the holding of final destination must be carried out under the control of the competent authority as long as the animals comprising the consignment are kept at the holding.
- (b) for ungulates for immediate slaughter, Article 11(2) shall apply.

Article 13

Conditions to be applied following the introduction into the Union of consignments of bees referred to in Article 7

- 1. Consignments of queen bees referred to in Article 7(3)(a) shall be conveyed without delay to the designated place of final destination where the hives shall be placed under the control of the competent authority and the queen bees transferred to new cages before being introduced to local colonies.
- 2. The cages, attendants, and other material that accompanied the queen bees from the third country of origin shall be sent to a laboratory designated by the competent authority for examination for the presence of:
- (a) the small hive beetle (Aethina tumida), their eggs or larvae;
- (b) signs of the Tropilaelaps mite (Tropilaelaps spp.).

After that laboratory examination, the cages, attendants and the material shall be destroyed.

3. Consignments of bumble bees (*Bombus spp.*) referred to in Article 7(3)(b) shall be conveyed without delay to the designated place of destination.

Those bumble bees may stay in the container in which they were introduced into the Union until the end of the lifespan of the colony.

That container and the material that accompanied the bumble bees from the third country of origin shall be destroyed at the end of the lifespan of the colony at the latest.

CHAPTER III

CONDITIONS FOR THE INTRODUCTION OF FRESH MEAT INTO THE UNION

Article 14

General conditions for the importation of fresh meat

Consignments of fresh meat intended for human consumption shall only be imported into the Union if they comply with the following conditions:

(a) they come from the third countries, territories or parts thereof listed in columns 1, 2 and 3 of the table in Part 1 of Annex II for which there is a model veterinary certificate corresponding to the consignment concerned listed in column 4 of the table in Part 1 of Annex II;

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- (b) they are presented at the border inspection post of introduction into the Union accompanied by the appropriate veterinary certificate, drawn up in accordance with the relevant model veterinary certificate set out in Part 2 of Annex II, taking into account the specific conditions indicated in column 6 of the table in Part 1 of that Annex, and completed and signed by an official veterinarian of the exporting third country;
- (c) they comply with the requirements set out in the veterinary certificate referred to in point (b), including:
 - (i) the supplementary guarantees laid down in that certificate, where indicated in column 5 of the table in Part 1 of Annex II;
 - (ii) any additional veterinary certification requirements that the Member State of destination may impose in accordance with Union veterinary legislation and which are included in the certificate.

Article 15

Conditions to be applied following the importation of unskinned carcases of wild cloven-hoofed game

In accordance with Article 8(2) of Council Directive 97/78/EC (¹), consignments of unskinned carcases of wild cloven-hoofed game for human consumption after further processing shall be conveyed without delay to the processing establishment of destination.

Article 16

Transit and storage of fresh meat

The introduction into the Union of consignments of fresh meat not intended for importation into the Union but destined for a third country either by immediate transit or after storage in the Union in accordance with Article 12(4) and Article 13 of Directive 97/78/EC, shall only be authorised if the consignments comply with the following conditions:

- (a) they come from the third countries, territories or parts thereof listed in columns 1, 2 and 3 of the table in Part 1 of Annex II, for which there is a model veterinary certificate corresponding to the consignment concerned listed in column 4 of the table in Part 1 of Annex II;
- (b) they comply with the specific animal health requirements for the consignment concerned, as set out in the relevant model veterinary certificate referred to in point (a);
- (c) they are accompanied by a veterinary certificate, drawn up in accordance with the model veterinary certificate set out in Annex III, and completed and signed by an official veterinarian of the exporting third country;

⁽¹⁾ OJ L 24, 30.1.1998, p. 9.

(d) they are certified as acceptable for transit, including for storage as appropriate, on the common veterinary entry document referred to in Article 2(1) of Commission Regulation (EC) No 136/2004 (¹), signed by the official veterinarian of the border inspection post of introduction into the Union.

Article 17

Derogation for transit through Latvia, Lithuania and Poland

- 1. By way of derogation from Article 16 the transit by road or by rail through the Union, between the designated border inspection posts in Latvia, Lithuania and Poland listed in Commission Decision 2009/821/EC (²), of consignments coming from and destined to Russia directly or via another third country shall be authorised provided that the following conditions are complied with:
- (a) the consignment is sealed with a serially numbered seal at the border inspection post of introduction into the Union by the veterinary services of the competent authority;
- (b) the documents accompanying the consignment and referred to in Article 7 of Directive 97/78/EC are stamped 'ONLY FOR TRANSIT TO RUSSIA VIA THE EU' on each page by the official veterinarian of the competent authority responsible for the border inspection post of introduction into the Union;
- (c) the procedural requirements provided for in Article 11 of Directive 97/78/EC are complied with;
- (d) the consignment is certified as acceptable for transit on the common veterinary entry document signed by the official veterinarian of the border inspection post of introduction into the Union.
- 2. Unloading or storage, as defined in Article 12(4) or in Article 13 of Directive 97/78/EC, of such consignments on Union territory shall not be allowed.
- 3. Regular audits shall be made by the competent authority to ensure that the number of consignments and the quantities of products leaving the Union territory matches the number and quantities entering.

CHAPTER IV

GENERAL, TRANSITIONAL AND FINAL PROVISIONS

Article 18

Certification

The veterinary certificates required by this Regulation shall be completed in accordance with the explanatory notes set out in Annex V.

⁽¹⁾ OJ L 21, 28.1.2004, p. 11.

⁽²⁾ OJ L 296, 12.11.2009, p. 1.

However, that requirement shall not preclude the use of electronic certification or of other agreed systems, harmonised at Union level.

Article 19

Transitional provisions

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For a transitional period those consignments of live animals, except bees coming from the State of Hawaii, and fresh meat intended for human consumption certified before 30 November 2010 in accordance with Decisions 79/542/EEC and 2003/881/EC may continue to be introduced into the Union until 31 May 2011.

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Article 20

Repeal

Decision 2003/881/EC is repealed.

Article 21

Entry into force

This Regulation shall enter into force on the twentieth day following that of its publication in the Official Journal of the European Union.

This Regulation shall be binding in its entirety and directly applicable in all Member States.

ANNEX I

UNGULATES

PART 1 List of third countries, territories or parts thereof (1)

ISO code and name of	Code of	Description of third country, territory or	Veterinary certi	Specific		
third country	Territory	part thereof	Model(s)	SG	conditions	
1 2		3	4	5	6	
	CA-0	Whole country	POR-X			
CA – Canada	Whole country, except the Okanagan Valley region of British Columbia described as follows: — From a point on the Canada/- United States border 120° 15′ longitude, 49° latitude — Northerly to a point 119° 35′ longitude, 50° 30′ latitude — North-easterly to a point 119° longitude, 50° 45′ latitude — Southerly to a point on the Canada/United States border 118° 15′longitude, 49° latitude		BOV-X, OVI-X, OVI-YRUM (*)		IVb IX	
CH – Switzerland	CH – Switzerland CH-0 Whole country		(**)			
CL – Chile	CL-0	Whole country	BOV-X, OVI-X, RUM			
		·	POR-X, SUI	В		
GL – Greenland	GL-0	Whole country	OVI-X, RUM		v	
HR – Croatia	HR-0	Whole country	BOV-X, BOV-Y, RUM, OVI-X, OVI-Y			
IS – Iceland	IS-0	Whole country	BOV-X, BOV-Y RUM, OVI-X, OVI-Y			
			POR-X, POR-Y	В		
ME – Montenegro	ME-0	Whole country			I	
MK – The former Yugoslav Republic of Macedonia (***)	MK-0	Whole country			I	

⁽¹⁾ Without prejudice to specific certification requirements provided for by any relevant agreement between the Union and third countries.

ISO code and name of	Code of Description of third country, territory or		Veterinary certi	Specific	
third country	Territory	part thereof	Model(s)	SG	conditions
1	2	3	4	5	6
NZ – New Zealand	NZ-0	Whole country	BOV-X, BOV-Y, RUM, POR-X, POR-Y OVI-X, OVI-Y		III V
PM – St Pierre and Miquelon	PM-0	Whole country	BOV-X, BOV-Y, RUM, OVI-X, OVI-Y CAM		
RS – Serbia (****)	RS-0	Whole country			I

(*) Exclusively for live animals other than animals belonging to the cervidae species.

(**) Certificates in accordance with the Agreement between the European Community and the Swiss Confederation on trade in agricultural products (OJ L 114, 30.4.2002, p. 132).

(***) The former Yugoslav Republic of Macedonia: the definitive nomenclature for this country will be agreed following current negotiations at UN level.

(****) Not including Kosovo which is at present under international administration pursuant to United Nations Security Council Resolution 1244 of 10 June 1999.

Specific Conditions (see footnotes in each certificate):

'1': for transit through the territory of a third country of live animals for immediate slaughter or live bovine animals for fattening which are consigned from a Member State and destined to another Member State in lorries which have been sealed with a serially numbered seal.

The seal number must be entered on the health certificate issued in accordance with the model laid down in Annex F to Directive 64/432/EEC (¹) for live bovine animals for slaughter and fattening and in accordance with Model I of Annex E to Directive 91/68/EEC (²) for ovine and caprine animals for slaughter.

In addition, the seal must be intact on arrival at the designated border inspection post of entry into the Union and the seal number recorded in the integrated computerised veterinary system of the Union (TRACES).

The certificate must be stamped at the exit point of the Union by the competent veterinary authority prior to transiting one or more third countries, with the following wording 'ONLY FOR TRANSIT BETWEEN DIFFERENT PARTS OF THE EUROPEAN UNION VIA THE FORMER YUGOSLAV REPUBLIC OF MACEDONIA/MONTENEGRO/SERBIA (*) (**)'.

Bovine animals for fattening must be transported directly to the holding of destination designated by the competent veterinary authority of destination. Those animals must not be moved from that holding unless for immediate slaughter.

(*) Delete country as applicable.

(**) Serbia does not include Kosovo which is at present under international administration pursuant to United Nations Security Council Resolution 1244 of 10 June 1999.

⁽¹⁾ OJ 121, 29.7.1964, p. 1977/64.

⁽²⁾ OJ L 46, 19.2.1991, p. 19.

▼C1

'II': territory recognised as having an official tuberculosis-free status for

the purposes of exports to the Union of live animals certified

according to the model of certificate BOV-X

'III': territory recognised as having an official brucellosis-free status for

the purposes of exports to the Union of live animals certified

according to the model of certificate BOV-X.

'IVa': territory recognised as having an official enzootic-bovine-

leukosis (EBL) free status for the purposes of exports to the Union of live animals certified according to the model of certificate

BOV -X.

▼<u>M2</u>

'IVb': recognised as having officially enzootic-bovine-leukosis (EBL)-free

herds equivalent to the requirements set out in Annex D to Directive 64/432/EEC for the purposes of exports to the Union of live animals

certified according to the model of certificate BOV-X.

▼<u>C1</u>

'V': territory recognised as having an official brucellosis-free status for

the purposes of exports to the Union of live animals certified

according to the model of certificate OVI-X.

'VI': Geographical constraints:

'VII': territory recognised as having an official tuberculosis-free status for

the purposes of exports to the Union of live animals certified

according to the model of certificate RUM.

'VIII': territory recognised as having an official brucellosis-free status for

the purposes of exports to the Union of live animals certified

according to the model of certificate RUM.

'IX': territory recognised as having an official Aujeszky's disease -free

status for the purposes of exports to the Union of live animals

certified according to the model of certificate POR-X.

PART 2

Models of Veterinary Certificates

Models:

'BOV-X': Model of veterinary certificate for domestic bovine animals

(including Bubalus and Bison species and their cross-breeds)

intended for breeding and/or production after importation.

'BOV-Y': Model of veterinary certificate for domestic bovine animals

(including Bubalus and Bison species and their cross-breeds)

intended for immediate slaughter after importation.

'OVI-X': Model of veterinary certificate for domestic ovine animals (Ovis

aries) and domestic caprine animals (Capra hircus) intended for

breeding and/or production after importation.

'OVI-Y': Model of veterinary certificate for domestic ovine animals (Ovis

aries) and domestic caprine animals (Capra hircus) intended for

immediate slaughter after importation.

'POR-X': Model of veterinary certificate for domestic porcine animals (Sus

scrofa) intended for breeding and/or production after importation;

'POR-Y': Model of veterinary certificate for domestic porcine animals (Sus

scrofa) intended for immediate slaughter after importation.

'RUM': Model of veterinary certificate for animals of the order Artiodactyla

(excluding bovine animals (including *Bubalus* and Bison species and their cross-breeds), *Ovis aries*, *Capra hircus*, *Suidae* and *Tayas-suidae*), and of the families *Rhinocerotidae* and *Elephantidae*.

'SUI': Model of veterinary certificate for non-domestic Suidae, Tayas-

suidae and Tapiridae.

'CAM': Model of specific attestation for animals imported from St Pierre and

Miquelon under the conditions provided for in Part 7 of Annex I.

SG (Supplementary guarantees):

'A': guarantees regarding Bluetongue and Epizootic-haemorrhagicdisease tests on animals certified according to the model of certificate BOV-X (point II.2.8 B), OVI-X (point II.2.6 D) and RUM (point II.2.6).

'B': guarantees regarding Swine-vesicular-disease and Classical-swine-fever tests on animals certified according to the model of certificate POR-X (point II.2.4 B) and SUI (point II.2.4 B).

'C': guarantees regarding Brucellosis test on animals certified according to the model of certificate POR-X (point II.2.4 C) and SUI (point

II.2.4 C).

▼<u>M2</u>

Model BOV-X

cou	NTRY	,	Veterinary certificate to EU			
	l.1.	Consignor	I.2. Certificate reference No I.2.a.			
		Name Address	I.3. Central competent authority			
			I.4. Local competent authority			
ent	1.5.	Tel. Consignee	1.6.			
signm		Name				
d con		Address				
pache		Postal code Tel.				
Part I: Details of dispached consignment	1.7.	Country of origin ISO code I.8. Region of origin Code origin	I.9. Country of destination ISO code destination Code destination			
l: Deta	1.11.	Place of origin	1.12.			
Part		Name Approval number Address				
	1.13.	Place of loading	I.14. Date of departure			
		Address Approval number				
	l.15.	Means of transport	I.16. Entry BIP in EU			
		Aeroplane				
		Identification Documentary references	1.17.			
	I.18.	Description of commodity	I.19. Commodity code (HS code) 01.02			
			I.20. Quantity			
	1.21.		I.22. Number of packages			
	1.23.	Seal/Container No	1.24.			
	1.25.	Commodities certified for:				
	r	Breeding	Fattening			
	1.26.		I.27. For import or admission into the EU			
	1.28.	Identification of the commodities	1			
		Species Breed Identification system (scientific name)	Identification number Age Sex			

▼<u>M2</u>

COL	UNTRY						Model BOV-X
	II.	Health	n informatio	n		II.a. Certificate reference number	II.b.
	II.1.	Public	Health Atte	estat	ion		
ion		I, the	undersigned	offici	al veterinarian, hereby certify, that the a	nimals described in this certificate:	
Part II: Certification		II.1.1.	brucellosis,	for the	ngs which have been free from any off ne past 30 days in the case of anthrax nals from holdings which did not satisfy	and for the past 6 months in the cas	
Part I		II.1.2.	have not re	ceive	d:		
			— any stilb	ene (or thyrostatic substances,		
					androgenic, gestagenic or β-agonist sub rective 96/22/EC),	stances for purposes other than there	speutic or zootechnic treatment (as
		II.1.3.	with regard	to bo	ovine spongiform encephalopathy (BSE):		
			(¹) (²) either	[(a)	the animals are identified by a permane herd of origin, and are not exposed bov of Regulation (EC) No 999/2001;		
				(b)	if there have been BSE indigenous cas which the ban on the feeding of rumin been effectively enforced or after the dat ban.]	ants with meat-and-bone meal and gr	eaves derived from ruminants had
		> ⁽¹⁾	³ (1)(3) or	[(a)	the animals are identified by a permane herd of origin, and are not exposed bov of Regulation (EC) No 999/2001; ◀		
				(b)	the animals were born after the date fr and greaves derived from ruminants h indigenous case if born after the date	nad been effectively enforced or after	
			(¹) (⁴) or	[(a)	the animals are identified by a permane herd of origin, and are not exposed bovi of Regulation (EC) No 999/2001;		
				(b)	the animals were born at least 2 years and-bone meal and greaves derived fro last BSE indigenous case if born after	m ruminants had been effectively enfo	
	II.2.	Anima	al Health att	estat	ion:		
		I, the	undersigned	offici	al veterinarian, hereby certify, that the a	nimals described above meet the follo	wing requirements:
		II.2.1.	they come	from	the territory with code:	(5) which, at the date of issuing this of	pertificate:
			(¹) either	[(a)	has been free for 24 months from foot-c fever, contagious bovine pleuropneumo months from vesicular stomatitis,]		
			(¹) or	[(a)	(i) has been free for 12 months from pneumonia, lumpy skin disease an stomatitis,	m rinderpest, bluetongue, Rift valley nd epizootic haemorrhagic disease,	
						and-mouth disease sincethat date, and authorised to expo	
			and	(b)	where during the last 12 months, no vidomestic cloven-hoofed animals vaccing		
		II.2.2.	they have re	emair	ned in the territory described under point	II.2.1. since birth, or for at least the la	ast 6 months before dispatch to the

COUNTRY Model BOV-X Ш Health information II.a. Certificate reference No II.b. II.2.3. they have remained since birth or at least 40 days before dispatch in the holding(s) of origin described under box (a) in and around which, in an area with a 150 km radius, there has been no case/outbreak of bluetongue and epizootic haemorrhagic disease during the previous 60 days, (b) in and around which, in an area with a 10 km radius, there has been no case/outbreak of the other diseases referred to in point II.2.1, during the previous 40 days: they are not animals to be killed under a national programme for the eradication of diseases, nor have they been vaccinated against the diseases referred to under point II.2.1.; 11.2.4. they come from herds that are not restricted under the national legislation pertaining to the eradication of tuberculosis, II.2.5. brucellosis and enzootic bovine leukosis; II.2.6. they come from herds recognised as officially tuberculosis-free (6); and (1) (7) either [come from a region which is recognised as officially tuberculosis-free (6);] [have been subjected to an intradermal tuberculin test (8) carried out with negative results within the past 30 days before dispatch to the Union;] (1) or (1) or [are less than 6 weeks old;] []].2.7. they have not been vaccinated against brucellosis and come from herds recognised as officially brucellosis-free (6); and (1) (7) either [come from a region which is recognised as officially brucellosis-free (6),] [have been subjected to at least one test for bovine brucellosis (8) carried out on samples taken within the (1) or past 30 days before dispatch to the Union.1 (1) or [are less than 12 months old,] [are castrated males of any age,] (1) either [II.2.8A. they come from herds included in an official system for the control of enzootic bovine leukosis, and in which there has been no evidence either clinical or as a result of a laboratory test of this disease during the past 2 years,] [II.2.8A. they come from herds recognised as officially enzootic-bovine-leukosis-free (6) (6a),] and $\binom{1}{7}$ either [come from a region which is recognised as officially enzootic-bovine-leukosis-free $\binom{6}{7}$] (1) or [have been subjected to an individual test for enzootic bovine leukosis (8) carried out with negative result on samples taken within the past 30 days before dispatch to the Union;] (1) or [are less than 12 months old;] $(^1)(^9)$ [II.2.8B. they have reacted negatively to a serological test for the detection of antibody for bluetongue and epizootic haemorrhagic within 10 days before export;] they are/were (1) dispatched from their holding(s) of origin, without passing through any market: (1) either [directly to the Union.] [to the officially authorised assembly centre described under box reference I.13. situated within the territory described under point II.2.1..] (1) or and, until dispatched to the Union: (a) they did not come in contact with other cloven-hoofed animals not complying with the health requirements as described

COUNTRY Model BOV-X

II. Health information II.a. Certificate reference No II.b.

- (b) they were not at any place where, or around which, within a 10 km radius, during the previous 30 days there has been a case/outbreak of any of the diseases referred to in point II.2.1.;
- II.2.10. any transport vehicles or containers in which they were loaded were cleaned and disinfected before loading with an officially authorised disinfectant:
- II.2.11. they were examined by an official veterinarian within 24 hours of loading and showed no clinical sign of disease;
- II.2.12. they have been loaded for dispatch to the Union on (dd/mm/yyyy) (10) in the means of transport described under box reference I.15. above that were cleaned and disinfected before loading with an officially authorised disinfectant and so constructed that faeces, urine, litter or fodder could not flow or fall out of the vehicle or container during transportation.

II.3. Animal transport attestation

I, the undersigned official veterinarian, hereby certify, that the animals described above have been treated before and at the time of loading in accordance with the relevant provisions of Regulation (EC) No 1/2005, in particular as regards watering and feeding, and they are fit for the intended transport.

(1) (11) [II.4. Specific requirements

- II.4.1. According to official information, no clinical or pathological evidence of infectious bovine rhinotracheitis (IBR) has been recorded in the holding(s) of origin referred to in box reference I.11., for the last 12 months;
- II.4.2. the animals referred to in box reference I.28.:
 - (a) have been isolated in accommodation approved by the competent authority for the last 30 days immediately prior to dispatch for export.
 - (b) have been subjected to a serological test for IBR on sera taken at least 21 days after entry into isolation, with negative results, and all animals in isolation have also given negative results to this test, and
 - (c) have not been vaccinated against IBR.]

Notes

This certificate is meant for domestic bovine animals (including *Bubalus* and *Bison* species and their cross-breeds) intended for breeding and/or production.

After importation the animals must be conveyed without delay to the holding of destination where they shall remain for a minimum period of 30 days before further movement outside the holding, except in the case of a dispatch to a slaughterhouse.

Part I

- Box reference I.8.: Provide the code of territory as appearing in Part 1 of Annex I to Regulation (EU) No 206/2010.
- Box reference I.13.: The assembly centre, if any, must fulfil the conditions for its approval, as laid down in Part 5 of Annex I to Regulation (EU) No 206/2010.
- Box reference I.15.: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor must inform the BIP of entry into the Union.
- Box reference I.23.: For containers or boxes, the container number and the seal number (if applicable) should be included.
- Box reference I.28.: Identification system: The animals must bear:
 - An individual number which permits tracing of their premises of origin. Specify the identification system (such as tag, tattoos, brand, chip, transponder),
 - An ear tag that includes the ISO code of the exporting country. The individual number must permit tracing of their premises of origin,

cou	NTRY		Model BOV-X					
II.	Health information	II.a. Certificate reference number	II.b.					
-	— Box reference I.28.: Species: Select amongst "Bos", "Bison" and "Bubalus" as appropriate.							
-	Box reference I.28.: Age: Date of birth (dd/mm/yy).							
-	Box reference I.28.: Sex (M = male, F = female, C = castrated).							
-	Box reference I.28.: Breed: select purebred, crossbreed.							
Par	t II:							
(1)	Keep as appropriate							
(2)	Only if the animals were born and continuously reared in a country 999/2001 as a country or region posing a negligible BSE risk and							
(3)	Only if the country or region of origin is categorised in accordance posing a controlled BSE risk and is listed as such in Decision 200		No 999/2001 as a country or region					
(4)	Only if the country or region of origin has not been categorised in categorised as a country or region with undetermined BSE risk an							
(5)	Code of the territory as it appears in Part 1 of Annex I to Regulat	ion (EU) No 206/2010						
(⁶)	Officially tuberculosis/brucellosis-free regions and herds as laid do regions and herds as laid down in Chapter I of Annex D to Direct		D; and enzootic-bovine-leukosis-free					
(^{6a})	Only for officially enzootic-bovine-leukosis-free herds recognised as equivalent to the requirements as laid down in Chapter I of Annex D to Directive 64/432/EEC for the purpose of exports to the EU of live animals according to the model certificate BOV-X from the territory that, in column 6 of Part 1 of Annex I to Regulation (EU) No 206/2010, appears with the entry "IVb" as regards enzootic bovine leukosis.							
(7)	Only for a territory that, in column 6 of Part 1 of Annex I to Regulat "III", as regards brucellosis, and/or "IVa" as regards enzootic boving the control of the column of		e entry "II", as regards tuberculosis,					
(8)	Tests carried out in accordance with the protocols that, for the dise $206/2010$.	ease concerned, are described in Part	6 of Annex I to Regulation (EU) No					
(⁹)	Supplementary guarantees to be provided when required in columentry "A".	nn 5 "SG" of Part 1 of Annex I to Reg	gulation (EU) No 206/2010, with the					
	Tests for bluetongue and for epizootic haemorrhagic disease in ac	ccordance with Part 6 of Annex I to Re	egulation (EU) No 206/2010.					
(10)	Date of loading. Imports of these animals shall not be allowed w exportation to the Union of the third country, territory or part ther measures have been adopted by the Union against imports of the	eof referred to in Boxes I.7 and I.8, o	or during a period where restrictive					
(11)	When required by the EU Member State of destination or Switzerla Agreement between the Community and the Swiss Confederation	and, in accordance with Decision 2004, on trade in agricultural products (OJ L	/558/EC and in accordance with the 114, 30.4.2002, p. 132).					
Offi	cial veterinarian							
	Name (in capital letters):	Qualification an	d title:					
	Date:	Signature:						
	Stamp:							

			del BOV-Y				
		UNTRY	Veterinary certificate to EU				
	l.1.	Consignor	I.2. Certificate reference number I.2.a.				
		Name	I.3. Central Competent Authority				
		Address	I.4. Local Competent Authority				
		Tel. No	1.4. Local Competent Authority				
ţ	1.5.	Consignee	1.6.				
nme		Name					
nsig		Address					
00 0		Postal code					
chec		Tel. No					
Part I: Details of dispatched consignment	1.7.	Country ISO I.8. Region Code of origin	I.9. Country of ISO I.10. Region of Code destination				
ls o	1.11	. Place of origin	I.12.				
l: Deta		Name Approval number Address					
Part		Name Approval number Address					
		Name Approval number Address					
	I.13	. Place of loading Address Approval number	I.14. Date of departure time of departure				
	I.15	. Means of transport Aeroplane Ship Railway wagon	I.16. Entry BIP in EU				
		Road vehicle Other Identification: Documentary references:	1.17.				
	1.18	. Description of commodity	I.19. Commodity code (HS code) 01.02				
			I.20. Quantity				
	I.21		I.22. Number of packages				
	1.23	B. Identification of container/seal number	1.24.				
	I.25. Commodities certified for: Slaughter						
	1.26	5.	I.27. For import or admission into EU				
	1.28	1					
		Species Breed Identification (Scientific name) system	Identification Age Sex number				

COUNTRY Model BOV-Y II. Health information II.a. Certificate reference number II.b. II.1. **Public Health Attestation** I, the undersigned official veterinarian, hereby certify, that the animals described in this certificate: come from holdings which have been free from any official prohibition on health grounds, for the last 42 days in the case of brucellosis, for the last 30 days in the case of anthrax, for the last six months in the case of rabies, and, have Part II: Certification not been in contact with animals from holdings which did not satisfy these conditions; II.1.2 have not received: any stilbene or thyrostatic substances, $oestrogenic, and rogenic, gestagenic \ or \ \beta\text{-} \ agonist \ substances \ for \ purposes \ other \ than \ the rapeutic \ or \ zootechnic$ treatment (as defined in Directive 96/22/EC). II.1.3 with regard to bovine spongiform encephalopathy (BSE): (1) (2) either [(a) the animals are identified by a permanent identification system enabling them to be traced back to the dam and herd of origin, and are not exposed bovine animals as described in Chapter C, part I, point 4) b) iv) of Annex II of Regulation (EC) No 999/2001; (b) if there have been BSE indigenous cases in the country concerned, 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 had been effectively enforced or after the date of birth of the last BSE indigenous case if born after the date of the feed ban.] (1) (3) or [(a) the animals are identified by a permanent identification system enabling them to be traced back to the dam and herd of origin, and are not exposed bovine animals as described in Chapter C, Part II, point (4)(b) iv) of Annex II of Regulation (EC) No 999/2001; (b) the animals were born after the date from which the ban on the feeding of ruminants with meatand-bone meal and greaves derived from ruminants had been effectively enforced or after the date of birth of the last BSE indigenous case if born after the date of the feed ban.] (1) (4) or [(a) the animals are identified by a permanent identification system enabling them to be traced back to the dam and herd of origin, and are not exposed bovine animals as described in Chapter C, Part II, point (4)(b)(iv) of Annex II of Regulation (EC) No 999/2001; (b) the animals were born at least two years after the date from which the ban on the feeding of ruminants with meat-and-bone meal and greaves derived from ruminants had been effectively enforced or after the date of birth of the last BSE indigenous case if born after the date of the feed ban.] 11.2. **Animal Health Attestation** I, the undersigned official veterinarian, hereby certify, that the animals described above meet the following requirements: 11.2.1 they come from the territory with code:(5) which, at the date of issuing this certificate: [(a) has been free for 24 months from foot-and-mouth disease, for 12 months from rinderpest, (1) either bluetongue, Rift valley fever, contagious bovine pleuropneumonia, lumpy skin disease and epizootic haemorrhagic disease, and for 6 months from vesicular stomatitis, and] [(a) (i) has been free for 12 months from rinderpest, bluetongue, Rift valley fever, contagious bovine pleuropneumonia, lumpy skin disease and epizootic haemorrhagic disease, and for (1) or 6 months from vesicular stomatitis, and has been considered free from foot-and-mouth disease since ... (dd/mm/yyyy), without having had cases/outbreaks from that date, and authorised to export these animals by Commission Regulation (EU) No, of .. (dd/mm/yyyy), and]

COUNTRY Model BOV-Y

II. Health information II.a. Certificate reference number II.b. (b) where during the last 12 months, no vaccination against these diseases has been carried out and imports of domestic cloven-hoofed animals vaccinated against these diseases are not permitted: 11.2.2 they have remained in the territory described under point II.2.1 since birth, or for at least the last three months before dispatch to the Union and without contact with imported cloven-hoofed animals for the last 30 days: 11.2.3 they have remained since birth or at least 40 days before dispatch in the holding(s) described under box reference I.11: (a) in and around which, in an area with a 150 km radius, there has been no case/outbreak of bluetongue and epizootic haemorrhagic disease during the previous 60 days, and (b) in and around which, in an area with a 10 km radius, there has been no case/outbreak of the other diseases referred to in point II.2.1 during the previous 40 days; they are not animals to be killed under a national programme for the eradication of diseases, nor have they been vaccinated against the diseases referred to in point II.2.1; II.2.5 they come from herds: (a) included in an official system for the control of enzootic bovine leukosis, and (b) that are not restricted under the national legislation regarding eradication of tuberculosis and brucellosis, and (c) recognised as officially tuberculosis free; (6) 11.2.6 they have not been vaccinated against brucellosis and they: (1) either [come from herds which are recognised as officially brucellosis free;] (6) (1) or [are castrated males of any age;] they are individually marked on at least two places on their hindquarters as to show that they are exclusively intended 11.2.7 for immediate slaughter; (7) 11.2.8 they are/were (1) dispatched from their holding(s) of origin, without passing through any market: (1) either [directly to the Union,] [to the officially authorised assembly centre described under box reference I.13 situated within the (1) or territory described under point II.2.1] and, until dispatched to the Union: (a) they did not come in contact with other cloven-hoofed animals not complying with the health requirements as described in this certificate, and (b) they were not at any place where, or around which within a 10 km radius, during the previous 30 days there has been a case/outbreak of any of the diseases referred to in point II.2.1: 11.2.9 any transport vehicles or containers in which they were loaded were cleaned and disinfected before loading with an officially authorised disinfectant: II.2.10 they were examined by an official veterinarian within 24 hours of loading and showed no clinical sign of disease; II.2.11 they have been loaded for dispatch to the Union on (dd/mm/yyyy) (8) in the means of transport described under box reference I.15 above that were cleaned and disinfected before loading with an officially authorised disinfectant and so constructed that faeces, urine, litter or fodder could not flow or fall out of the vehicle or container during transportation.

II.3. **Animal transport attestation**

I, the undersigned official veterinarian, hereby certify, that the animals described above have been treated before and at the time of loading in accordance with the relevant provisions of Regulation (EC) No 1/2005, in particular as regards watering and feeding, and they are fit for the intended transport.

COUNTRY Model BOV-Y

II.	Health information	II.a. Certificate reference number	II.b.

Notes

This certificate is meant for live bovine animals (including *Bubalus* and *Bison* species and their cross-breeds) intended for immediate slaughter.

After importation the animals must be conveyed without delay to the slaughterhouse of destination to be slaughtered within five working days.

Part I

- Box reference I.8: Provide the code of territory as appearing in Part 1 of Annex I to Regulation (EU) No 206/2010.
- Box reference I.13: The assembly centre, if any, must fulfil the conditions for its approval, as laid down in Part 5 of Annex I to Regulation (EU) No 206/2010.
- Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor must inform the BIP of entry into the Union.
- Box reference I.23: For containers or boxes, the container number and the seal number (if applicable) should be included.
- Box reference I.28: Identification system: the animals must bear:
 - An individual number which permits tracing of their premises of origin. Specify the identification system (such as tag, tattoos, brand, chip, transponder).
 - An ear tag that includes the ISO code of the exporting country. The individual number must permit tracing of their premises of origin.
- Box reference I.28: Species: Select amongst 'Bos', 'Bison' and 'Bubalus' as appropriate.
- Box reference I.28: Age: Date of birth (dd/mm/yy).
- Box reference I.28: Sex (M = male, F = female, C = castrated).

Part II:

- (1) Keep as appropriate.
- (2) Only if the animals were born and continuously reared in a country or region categorised in accordance with Article 5(2) of Regulation (EC) No 999/2001 as a country or region posing a negligible BSE risk and listed as such in Decision 2007/453/EC.
- (3) Only if the country or region of origin is categorised in accordance with Article 5(2) of Regulation (EC) No 999/2001 as a country or region posing a controlled BSE risk and is listed as such in Decision 2007/453/EC.
- (4) Only if the country or region of origin has not been categorised in accordance with Article 5(2) of Regulation (EC) No 999/2001 or has been categorised as a country or region with undetermined BSE risk and is listed as such in Decision 2007/453/EC.
- (5) Code of the territory as it appears in Part 1 of Annex I to Regulation (EU) No 206/2010.
- (6) Officially tuberculosis/brucellosis free regions and herds as laid down in Annex A to Directive 64/432/EEC.
- (7) This mark shall take the form of 'L' having 13 cm in the left side and 7 cm in the bottom side with 1 cm of strength in both lines. It shall be applied using the technique known as 'freeze-branding'.
- (a) Date of loading. Imports of these animals shall not be allowed when the animals were loaded either prior to the date of authorisation for exportation to the Union of the third country, territory or part thereof referred to in boxes I.7 and I.8, or during a period where restrictive measures have been adopted by the Union against imports of these animals from this third country, territory or part thereof.

l									
	Official veterinarian								
	Name (in capital letters):	Qualification and title:							
	Date:	Signature:							
	Stamp:								
	Stamp:								

			Mode	OVI-X				
		UNTRY					Veterinary cer	rtificate to EU
	l.1.	Consignor		I.2. Certificate re	eference	number	I.2.a.	
		Name		I.3. Central Com	petent A	uthority		
		Address		I.4. Local Comp	etent Aut	hority		
		Tel. No		•				
ent	1.5.	Consignee		I.6.				
gum		Name						
nsić		Address						
o p		Postal code						
tche		Tel. No						
Part I: Details of dispatched consignment	1.7.	Country ISO I.8. R of origin code of	legion Code f origin	I.9. Country of destination		SO ode	I.10. Region of destination	Code
ails o	1.11	. Place of origin		I.12.				
t I: Deta		Name Appro Address	val number					
Pari		Name Appro Address	val number			/		
		Name Appro Address	val number					
	I.13	. Place of loading Address Appro	val number	I.14. Date of departure time of departure				
	I.15	. Means of transport Aeroplane Ship Ship	Railway wagon 🔲	I.16. Entry BIP in EU				
		Road vehicle Other Identification: Documentary references:		1.17.				
	1.18	. Description of commodity		I.19. Commodity code (HS code)				
						I.20. Q	uantity	
	1.21					I.22. N	umber of package	es
	1.23	3. Identification of container/seal numb	per			1.24.		
	1.25	6. Commodities certified for:						
		Breeding		Fatte	ening 🗌			
	1.26	5.	I.27. For import o	r admissi	on into E	EU		
	1.28	3. Identification of the commodities						
		Species Breed (Scientific name)	Identification system	Identification number	1	Ag	е	Sex

COUNTRY Model OVI-X II. Health information II.a. Certificate reference number II.b. II.1. **Public Health Attestation** I, the undersigned official veterinarian, hereby certify, that the animals described in this certificate: come from holdings which have been free from any official prohibition on health grounds, for the last 42 days in the case of brucellosis, for the last 30 days in the case of anthrax, for the last six months in the case of rabies, and, have Part II: Certification not been in contact with animals from holdings which did not satisfy these conditions; II.1.2 have not received: any stilbene or thyrostatic substances, oestrogenic, androgenic, gestagenic or β - agonist substances for purposes other than therapeutic or zootechnic treatment (as defined in Directive 96/22/EC). 11.2. **Animal Health attestation** I, the undersigned official veterinarian, hereby certify, that the animals described above meet the following requirements: they come from the territory with code:(2) which, at the date of issuing this certificate: (1) either [(a) has been free for 24 months from foot-and-mouth disease, for 12 months from rinderpest, bluetongue, Rift valley fever, peste des petits ruminants, sheep pox and goat pox, contagious caprine pleuropneumonia and epizootic haemorrhagic disease and for 6 months from vesicular stomatitis, and] (1) or [(a) (i) has been free for 12 months from rinderpest, bluetongue, Rift valley fever, peste des petits ruminants, sheep pox and goat pox, contagious caprine pleuro-pneumonia and epizootic haemorrhagic disease, and for 6 months from vesicular stomatitis, and (ii) has been considered free from foot-and-mouth disease, since .. (dd/mm/yyyy), without having had cases/outbreaks from that date, and authorised to export these animals by Commission Regulation (EU) No, of (dd/mm/yyyy), and] (b) where during the last 12 months, no vaccination against these diseases has been carried out and imports of domestic cloven-hoofed animals vaccinated against these diseases are not they have remained in the territory described under point II.2.1 since birth, or for at least the last six months before 11.2.2 dispatch to the Union and without contact with imported cloven-hoofed animals for the last 30 days; 11.2.3 they have remained since birth or at least 40 days in the holding(s) described under box reference I.11 before (a) in and around which, in an area with a 150 km radius, there has been no case/outbreak of bluetongue and epizootic haemorrhagic disease during the previous 60 days, and (b) in and around which, in an area with a 10 km radius, there has been no case/outbreak of the other diseases referred to in point II.2.1 during the previous 40 days; II.2.4 according to my knowledge and to the written declaration made by the owner, the animals: (a) do not come from holdings, and have not been in contact with animals of a holding, in which the following diseases have been clinically detected: (i) contagious agalactia of sheep or goats (Mycoplasma agalactiae, Mycoplasma capricolum, Mycoplasma mycoides var. mycoides large colony), within the last six months, (ii) paratuberculosis and caseous lymphadenitis, within the last 12 months,

(iii) pulmonary adenomatosis, within the last three years, and

(iv) Maedi/Visna or caprine viral arthritis/encephalitis:

COUNTRY Model OVI-X

II. Health information			II.a. Certificate reference number	II.b.	
	(¹) eith	er	within the last three years,]		
	(¹) or		within the last 12 months, and all the infected a animals subsequently reacted negatively to apart,]		
	(b) are include	ed in an of	icial system for notification of these diseases,	and	
	(c) have been export;	free from	clinical or other evidence of tuberculosis and	I brucellosis during the three years prior to	
II.2.5			be killed under a national programme for the elseases referred to in point II.2.1;	eradication of diseases, nor have they been	
II.2.6 A	they originate:				
	(¹) (³) either	[from the	e territory described under box reference I.8 sis-free;]	, which has been recognised as officially	
	(¹) or	[from the melitens	holding(s) described under box reference I.1 is):	1, where, in respect of brucellosis (Brucella	
			usceptible animals have been free from clinic nonths,	cal or any signs of this disease for the last	
			presentative number of the domestic ovine and submitted each year to a serological test, (4)	d caprine animals over an age of six months	
			omestic ovine or caprine animals have not be vaccinated with Rev. 1 vaccine more than two		
(dd/ı			ast two tests (°), separated by an interval of a mm/yyyy) and on(dd/mm/yyyy) on a nonths of age gave negative results, and]		
	(¹) or		nestic ovine or caprine animals under the age of 7 months are vaccinated against this ease with Rev. 1 vaccine;		
		(d) the I	last two tests (6), separated by an interval of at least six months, carried out:		
			on(dd/mm/yyyy) and dall non-vaccinated domestic ovine and caprine		
			on(dd/mm/yyyy) and on vaccinated domestic ovine and caprine animals		
		gave	negative results, and]		
			e are only domestic ovine and caprine animals irements;]	that fulfil at least the above conditions and	
(¹) [II.2.6 B	contagious epi	didymitis (ave been kept continuously during the previor Brucella ovis) has been diagnosed in the last 1 ays a complement fixation test to detect conta	2 months and, these rams have undergone	
II.2.6 C	In respect of so	crapie			
(¹) (²) [II.2.6.C.1	(¹) (²) [II.2.6.C.1 if they are destined for a Member State which benefits, for all or part of its territory, from the provisions laid down in point (b) or (c) of Chapter A(I) of Annex VIII to Regulation (EC) No 999/2001, the animals comply with the guarantees provided for in the programmes referred to in those points and the animals comply with the guarantees requested by the EU Member States of destination regarding scrapie, and]				
	either				
(¹) [II.2.6.C.2	are animals in		production born in and continuously reared of	on holdings in which a case of scrapie has	

COUNTRY Model OVI-X

II.	Health	information		II.a. Certificate reference numl	ber	II.b.
(¹) (8) or [II	I.2.6.C.2			t continuously since birth or for uirements for at least three year		years on a holding or holdings which have
		— they are sul	bject to re	gular official veterinary checks,		
		— the animals	are ident	ified in conformity with Union leg	islation,	
		— no case of	scrapie ha	s been confirmed;		
		the framew	ork of a di in accord	sease eradication campaign or s ance with the laboratory metho	slaughtered fo	on the holdings (except the animals killed in r human consumption) have been examined in point 3.2(b) of Chapter C of Annex X to
			ave been			ovine animals of the ARR/ARR prion protein rom holdings which complies with the above
(¹) or [II.2	.6.C.2	they are dome 2002/1003/EC;		animals of the ARR/ARR price	on protein ge	notype, as defined in Annex I to Decision
(¹) (⁹) [II	I.2.6 D	haemorrhagic-c quarantine perio	disease, o	arried out on two occasions on	samples of b	on of antibody for bluetongue and epizootic- lood taken at the beginning of the isolation (dd/mm/yyyy) and onys of export;]
	II.2.7	they are/were (1) dispatch	ned from their holding(s) of origin	n, without pass	sing through any market,
		(1) either	[directly	o the Union,]		
		(¹) or		ficially authorised assembly cen described under point II.2.1]	tre described	under box reference I.13 situated within the
		and, until dispa	tched to t	ne Union:		
		(a) they did no described in			ed animals no	t complying with the health requirements as
				place where, or around which with the control of the diseases referred		adius, during the previous 30 days there has 1.1;
	II.2.8	any transport ve officially author			aded were cle	eaned and disinfected before loading with an
	II.2.9	they were exam	ined by a	n official veterinarian within 24 h	ours of loadin	g and showed no clinical sign of disease;
	II.2.10	transport description	ribed und ised disin	er box reference I.15 above that	at were clean	(dd/mm/yyyy) (10) in the means of ed and disinfected before loading with an litter or fodder could not flow or fall out of the
II.3 .	Anima	transport attes	station			
	at the t	ime of loading in	n accorda		s of Regulation	ribed above have been treated before and on (EC) No 1/2005, in particular as regards

Notes

This certificate is meant for live domestic ovine animals (Ovis aries) and domestic caprine animals (Capra hircus) intended for breeding or production.

COUNTRY Model OVI-X

II. Health information II.a. Certificate reference number II.b.

After importation the animals must be conveyed without delay to the holding of destination where they shall remain for a minimum period of 30 days before further movement outside the holding, except in the case of a dispatch to a slaughterhouse.

Part I:

- Box reference I.8: Provide the code of territory as appearing in Part 1 of Annex I to Regulation (EU) No 206/2010.
- Box reference I.13: The assembly centre, if any, must fulfil the conditions for its approval, as laid down in Part 5 of Annex I to Regulation (EU) No 206/2010.
- Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor must inform the BIP of entry into the Union.
- Box reference I.19: Use the appropriate HS code: 01.04.10 or 01.04.20.
- Box reference I.23: For containers or boxes, the container number and the seal number (if applicable) should be included.
- Box reference I.28: Identification system: The animals must bear:
 - An individual number which permits tracing of their premises of origin. Specify the identification system (such as tag, tattoos, brand, chip, transponder) and the anatomic place used in the animal.
 - An ear tag that includes the ISO code of the exporting country. The individual number must permit tracing of their premises of origin.
- Box reference I.28: Species: Select amongst 'Ovis aries' and 'Capra hircus' as appropriate.
- Box reference I.28: Age: (months).
- Box reference I.28: Sex (M = male, F = female, C = castrated).

Part II:

- (1) Keep as appropriate.
- (2) Code of the territory as it appears in Part 1 of Annex I to Regulation (EU) No 206/2010.
- (3) Only for a territory appearing with the entry 'V' in column 6 of Part 1 of Annex I to Regulation (EU) No 206/2010.
- (4) The representative number of animals to be tested for brucellosis must, for each holding, consist of:
 - all non-castrated male animals, which have not been vaccinated against brucellosis, over six months old,
 - all non-castrated male animals, which have been vaccinated against brucellosis, over 18 months old,
 - all animals brought onto the holding since the previous tests, and
 - $-25\,\%$ of females which are sexually mature, within a minimum of 50 females.
- (5) This must be completed when the destination is a Member State or part of a Member State laid down in one of the Annexes of Decision 93/52/EEC.
- (6) In accordance with Part 6 of Annex I to Regulation (EU) No 206/2010.
 - Where more than one holding of origin is involved the date of the most recent test on each holding must be clearly indicated.
- (7) Guarantees in relation to a programme of control of scrapie, as requested by the EU Member State of destination, in application of Article 15 and Annex IX, Chapter E of Regulation (EC) No 999/2001.
- (8) In the case of animals intended, exclusively, for breeding purposes.
- (9) Supplementary guarantees to be provided when required in column 5 'SG' of Part 1 of Annex I to Regulation (EU) No 206/2010, with the entry 'A'. Tests for Bluetongue and for Epizootic-haemorrhagic-disease in accordance with Part 6 of Annex I to Regulation (EU) No 206/2010.
- (10) Date of loading. Imports of these animals shall not be allowed when the animals were loaded either prior to the date of authorisation for exportation to the Union of the third country, territory or part thereof referred to in boxes I.7 and I.8, or during a period where restrictive measures have been adopted by the Union against imports of these animals from this third country, territory or part thereof.

COUNTRY		
II. Health information	II.a. Certificate reference number	II.b.
Official veterinarian		
Name (in capital letters):	Qualification	and title:
Date:	Signature:	
Stamp:		

tched consignment	.1. Consignor Name Address Tel. No .5. Consignee Name	1.2. Certificate reference number 1.2.a. 1.3. Central Competent Authority 1.4. Local Competent Authority 1.6.	
	Name Address Tel. No .5. Consignee	I.3. Central Competent Authority I.4. Local Competent Authority	
atched consignment	Address Tel. No .5. Consignee	I.4. Local Competent Authority	
atched consignment	Tel. No .5. Consignee		
atched consignment	.5. Consignee	16	
atched consignmen			
atched consign			
atched cons	Address		
atched	Postal code		
atc	Tel. No		
disp .	.7. Country ISO I.8. Region Code of origin	I.9. Country of ISO I.10. Region of Code destination	
18 1.	.11. Place of origin	I.12.	
I: Deta	Name Approval number Address		
Part	Name Approval number Address		
	Name Approval number Address		
1.	.13. Place of loading	I.14. Date of departure time of departure	
	Address Approval number		
1.	.15. Means of transport Aeroplane Ship Railway wagon	I.16. Entry BIP in EU	
	Road vehicle Other	I.17.	
	Identification: Documentary references:		
1.	.18. Description of commodity	I.19. Commodity code (HS code)	
		I.20. Quantity	
I.	.21.	I.22. Number of packages	
I.	.23. Identification of container/seal number	1.24.	
I.25. Commodities certified for: Slaughter			
1.	.26.	I.27. For import or admission into EU	
1.	.28. Identification of the commodities		
	Species Breed Identification (Scientific name) system	Identification Age Sex number	

COUNTRY Model OVI-Y II. Health information II.a. Certificate reference number II.b. II.1. **Public Health Attestation** I, the undersigned official veterinarian, hereby certify, that the animals described in this certificate: come from holdings which have been free from any official prohibition on health grounds, for the last 42 days in the case of brucellosis, for the last 30 days in the case of anthrax, for the last six months in the case of rabies, and, have Part II: Certification not been in contact with animals from holdings which did not satisfy these conditions; II.1.2 have not received: any stilbene or thyrostatic substances, oestrogenic, androgenic, gestagenic or β - agonist substances for purposes other than therapeutic or zootechnic treatment (as defined in Directive 96/22/EC). II.2. **Animal Health attestation** I, the undersigned official veterinarian, hereby certify, that the animals described above meet the following requirements: II.2.1 they come from the territory with code:(1) which, at the date of issuing this certificate: (2) either [(a) has been free for 24 months from foot-and-mouth disease, for 12 months from rinderpest, bluetongue, Rift valley fever, peste des petits ruminants, sheep pox and goat pox, contagious caprine pleuro-pneumonia and epizootic haemorrhagic disease and for 6 months from vesicular stomatitis, and] [(a) (i) has been free for 12 months from rinderpest, bluetongue, Rift valley fever, peste des petits (2) or ruminants, sheep pox and goat pox, contagious caprine pleuro-pneumonia and epizootic haemorrhagic disease, and for 6 months from vesicular stomatitis, and (ii) has been considered free from foot-and-mouth disease, since (dd/mm/yyyy), without having had cases/outbreaks from that date, and authorised to export these animals by Commission Regulation (EU) No, of ... (dd/mm/yyyy), and] (b) where during the last 12 months, no vaccination against these diseases has been carried out and imports of domestic cloven-hoofed animals vaccinated against these diseases are not permitted: they have remained in the territory described under point II.2.1 since birth, or for at least the last three months before 11.2.2 dispatch to the Union and without contact with imported cloven-hoofed animals for the last 30 days: II.2.3 they have remained since birth or at least 40 days before dispatch in the holding(s) described under box reference I.11: (a) in and around which in an area with a 150 km radius there has been no case/outbreak of bluetongue and epizootic haemorrhagic disease during the previous 60 days, and (b) in and around which, in an area with a 10 km radius, there has been no case/outbreak of the other diseases referred to in point II.2.1 during the previous 40 days; 11.2.4 they are not animals to be killed under a national programme for the eradication of diseases, nor have they been vaccinated against the diseases referred to in point II.2.1; II.2.5 they are/were (2) dispatched from their holding(s) of origin, without passing through any market, (2) either [directly to the Union]

territory described under point II.2.1,]

[to the officially authorised assembly centre described under box reference I.13 situated within the

(2) or

COUNTRY Model OVI-Y

II.	II. Health information			II.a. Certificate reference number	II.b.		
		and, until disp	atched to t	he Union:			
				contact with other cloven-hoofed animals not ificate, and	t complying with the health requirements as		
				place where, or around which within a 10 km ra k of any of the diseases referred to in point II.2			
	II.2.6	in respect of s	crapie:				
		(2) (3)	provisior comply v	are destined for a Member State which bene is laid down in point (b) or (c) of Chapter A(l) of with the guarantees provided for in the program 2 of Regulation (EC) 546/2006, and]	Annex VIII to Regulation (EC) No 999/2001,		
		(²) either	[were bo	orn in and continuously reared on holdings in ed;]	n which a case of scrapie has never been		
	Decision			mestic ovine animals of the ARR/ARR prion protein genotype as defined in Annex I to in 2002/1003/EC, coming from a holding where no case of scrapie has been reported in the nonths;]			
	II.2.7	any transport officially author		containers in which they were loaded were cle fectant;	eaned and disinfected before loading with an		
	II.2.8	they were exa	mined by a	n official veterinarian within 24 hours of loading	g and showed no clinical sign of disease;		
	II.2.9	transport desc officially autho	cribed und rised disin	for dispatch to the Union oner box reference I.15 above that were clean fectant and so constructed that faeces, urine, I g transportation.	ed and disinfected before loading with an		

II.3. Animal transport attestation

I, the undersigned official veterinarian, hereby certify, that the animals described above have been treated before and at the time of loading in accordance with the relevant provisions of Regulation (EC) No 1/2005, in particular as regards watering and feeding, and they are fit for the intended transport.

Notes

This certificate is meant for live domestic ovine animals (*Ovis aries*) and domestic caprine animals (*Capra hircus*) intended for immediate slaughter after importation.

After importation the animals must be conveyed without delay to the slaughterhouse of destination to be slaughtered within five working days.

COUNTRY	Model OVI-Y
COUNTRY	Model OVI-Y

II.	Health information	II.a. Certificate reference number	II.b.			
Pa	rt I:					
_	Box reference I.8: Provide the code of te	erritory as appearing in Part 1 of Annex I to Reg	gulation (EU) No 206/2010.			
_	- Box reference I.13: The assembly centre, if any, must fulfil the conditions for its approval, as laid down in Part 5 of Annex I to Regulation (EU) No 206/2010.					
_		r (railway wagons or container and lorries), flig ding, the consignor must inform the BIP of ent				
_	Box reference I.19: Use the appropriate	HS code: 01.04.10 or 01.04.20.				
_	Box reference I.23: For containers or bo	xes, the container number and the seal numbe	er (if applicable) should be included.			
_	Box reference I.28: Identification system	:The animals must bear:				
	 An individual number which permits brand, chip, transponder) and the ar 	tracing of their premises of origin. Specify the natomic place used in the animal.	e identification system (such as tag, tattoos,			
	 An ear tag that includes the ISO coo origin. 	de of the exporting country. The individual num	ber must permit tracing of their premises of			
_	Box reference I.28: Species: Select amo	ngst 'Ovis aries' and 'Capra hircus' as appropr	iate.			
_	Box reference I.28: Age: months.					
_	Box reference I.28: Sex (M = male, F = f	emale, C = castrated).				
Pa	rt II:					
(¹)	Code of the territory as it appears in Par	t 1 of Annex I to Regulation (EU) No 206/2010				
(2)	Keep as appropriate.					
(3)	Guarantees in relation to a programme of Article 15 and Chapter E of Annex IX to	of control of scrapie, as requested by the EU M Regulation (EC) No 999/2001.	ember State of destination, in application of			
(4)	for exportation to the Union of the third	s shall not be allowed when the animals were lo country, territory or part thereof referred to in d by the Union against imports of these anim	boxes I.7 and I.8, or during a period where			

Official	veterinarian	

Name (in capital letters):	Qualification and title

Date: Signature:

Stamp:

	Model POR-X							
		UNTRY		T			Veterinary cer	tificate to EU
	I.1.	Consignor		I.2. Certific	ate reference i	number	1.2.a.	
		Name		I.3. Central	Competent Au	uthority		
		Address		I4 Local C	Competent Auti	hority		
		Tel. No			- The term of the			
ent	1.5.	Consignee		1.6.				
n us		Name						
nsić		Address						
o p		Postal code						
tche		Tel. No						
Part I: Details of dispatched consignment	1.7.	Country ISO code	I.8. Region Code of origin	I.9. Country destina		SO ode	I.10. Region of destination	Code
ils o	1.11	. Place of origin		I.12.				
l: Deta		Name Address	Approval number					
Part		Name Address	Approval number					
		Name Address	Approval number					
	I.13	. Place of loading Address	Approval number	I.14. Date of departure time of departure				
	I.15	. Means of transport Aeroplane Sh	ip 🗌 Railway wagon 🔲	I.16. Entry BIP in EU				
		Road vehicle Oth Identification: Documentary references:	er 🗌	1.17.				
	I.18	. Description of commodity		I.19. Commodity code (HS code) 01.03				01.03
						I.20. Q	uantity	
	I.21			I.22. Number of packages			es	
	1.23	3. Identification of container/s	eal number	1.24.				
	1.25	5. Commodities certified for:			L			
		Breeding			Fattening			
	1.26	5.	I.27. For import or admission into EU					
	1.28	B. Identification of the commo	dities					
		Species (Scientific name)	Identification system	Identification number	n	Age	е	Sex

COUNTRY Model POR-X II. Health information II.a. Certificate reference number II.b. II.1. **Public Health Attestation** I, the undersigned official veterinarian, hereby certify, that the animals described in this certificate: come from holdings which have been free from any official prohibition on health grounds, for the last 42 days in the case of brucellosis, for the last 30 days in the case of anthrax and for the past six months in the case of rabies and, Part II: Certification the animals have not been in contact with animals from holdings which did not satisfy these conditions; II.1.2 have not received: any stilbene or thyrostatic substances, oestrogenic, androgenic, gestagenic or β - agonist substances for purposes other than therapeutic or zootechnic treatment (as defined in Directive 96/22/EC). II.2. **Animal Health attestation** I, the undersigned official veterinarian, hereby certify, that the animals described above meet the following requirements: II.2.1 they come from the territory with code:(1) which, at the date of issuing this certificate: (2) either [(a) has been free for 24 months from foot-and-mouth disease, for 12 months from rinderpest, African swine fever, classical swine fever, swine vesicular disease and vesicular exanthema, and for 6 months from vesicular stomatitis, and] $\hbox{\it [(a) (i)} \quad \hbox{has been free [for 24 months from foot-and-mouth disease] (2), for 12 months from rinderpest,}$ (2) or African swine fever, vesicular exanthema, [classical swine fever] (2) and [swine vesicular disease] (2), and for 6 months from vesicular stomatitis, and (ii) has been considered free from [foot-and-mouth disease] (2), [classical swine fever] (2) and [swine vesicular disease] (2), since (dd/mm/yyyy), without having had cases/outbreaks from that date, and authorised to export these animals by Commission (b) where during the last 12 months, no vaccination against these diseases has been carried out and imports of domestic cloven-hoofed animals vaccinated against these diseases are not II.2.2 they have remained in the territory described under point II.2.1 since birth, or for at least the last six months before dispatch to the Union and without contact with imported cloven-hoofed animals for the last 30 days; 11.2.3 they have remained in the holding(s) described under box reference I.11 since birth, or for at least 40 days prior to dispatch, and, during this period, in the holding(s) and in an area with a 10 km radius around the holding(s) of origin, there has been no case/outbreak of the diseases referred to in point II.2.1; II.2.4 A they are not animals to be killed under a national programme for the eradication of diseases, nor have they been vaccinated against the diseases referred to in point II.2.1; (2) (3) [II.2.4 B they have been subjected within the past 30 days to a test for swine vesicular disease antibodies and a test for classical swine fever antibodies with negative results in both cases]; (2) (4) [II.2.4 C they have been subjected within the past 30 days to a buffered Brucella antigen test for porcine brucellosis with negative results]: II.2.5 they come from herds which are not restricted under the national brucellosis eradication programme; II.2.6 they are/were (2) dispatched from their holding(s) of origin, without passing through any market, (2) either [directly to the Union,]

territory described under point II.2.1,]

(2) or

[to the officially authorised assembly centre described under box reference I.13 situated within the

COUNTE	COUNTRY Model POR-						
II.		info	rmation	II.a. Certificate refer	ence number	II.b.	_
	II.2.7 II.2.8 II.2.9	(a) (b) any off the	described in this cert they were not at any been a case/outbrea y transport vehicles or icially authorised disin by were examined by a ey have been loaded	contact with other clo ifficate, and place where, or around k of any of the disease containers in which the fectant; in official veterinarian veter of the L	d which within a 10 km s referred to in point II.3 ey were loaded were clayithin 24 hours of loadir	ot complying with the health requirements a radius, during the previous 40 days there has 2.1; eaned and disinfected before loading with a rig and showed no clinical sign of disease;	ıs .n
II.3.	I, the u	vel II tra unde time	hicle or container during the state of the s	ng transportation. narian, hereby certify,	that the animals desc provisions of Regulati	litter or fodder could not flow or fall out of th cribed above have been treated before an on (EC) No 1/2005, in particular as regard	d
(²) (⁶) [II.4	Specif		equirements				
	[11.4.1			•	eferred to in box referer		
	II.4.2	rec		months in the holding		al evidence of Aujeszky's disease has bee in box reference I.11, and in those holding	
	11.4.3	the	e animals referred to in	box reference I.28:			
		(a)				the holding(s) of origin referred to in bo he last 3 months and in others of equivaler	
		(b)			roved by the compete rindirect contact with o	nt authority for the last 30 days immediatel ther Suidae animals,	y
		(c)				body (7) on sera taken at least 21 days afte n have also given negative results to this tes	
		(d)			's disease and have not during the previous 12 i	t been in contact with vaccinated animals an months.]	d

Notes

This certificate is meant for live domestic porcine animals (Sus scrofa) intended for breeding or production.

After importation the animals must be conveyed without delay to the holding of destination where they shall remain for a minimum period of 30 days before further movement outside the holding, except in the case of a dispatch to a slaughterhouse.

(2) (8) [II.4.4 (further requirements and/or tests)

COUNTRY Model POR-X

II.	Health information	II.a. Certificate reference number	II.b.

Part I:

- Box reference I.8: Provide the code of territory as appearing in Part 1 of Annex I to Regulation (EU) No 206/2010.
- Box reference I.13: The assembly centre, if any, must fulfil the conditions for its approval, as laid down in Part 5 of Annex I to Regulation (EU) No 206/2010.
- Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor must inform the BIP of entry into the Union.
- Box reference I.23: For containers or boxes, the container number and the seal number (if applicable) should be included.
- Box reference I.28: Identification system: the animals must bear:
 - An individual number which permits tracing of their premises of origin. Specify the identification system (such as tag, tattoos, brand, chip, transponder).
 - An ear tag that includes the ISO code of the exporting country. The individual number must permit tracing of their premises of origin.
- Box reference I.28: Age: months.
- Box reference I.28: Sex (M = male, F = female, C = castrated).

Part II:

- (1) Code of the territory as it appears in Part 1 of Annex I to Regulation (EU) No 206/2010.
- (2) Keep as appropriate.
- (3) Supplementary guarantees to be provided when required in column 5 'SG' of Part 1 of Annex I to Regulation (EU) No 206/2010, with the entry 'B'.
- (4) Supplementary guarantees to be provided when required in column 5 'SG' of Part 1 of Annex I to Regulation (EU) No 206/2010, with the entry 'C'.
- (5) Date of loading. Imports of these animals shall not be allowed when the animals were loaded either prior to the date of authorisation for exportation to the Union of the third country, territory or part thereof referred to in boxes I.7 and I.8, or during a period where restrictive measures have been adopted by the Union against imports of these animals from this third country, territory or part thereof.
- (*) When required by the EU Member State of destination or Switzerland, in accordance with Decision 2008/185/EC and the Agreement between the Community and the Swiss Confederation on trade in agricultural products (OJ L 114, 30.4.2002, p. 132) except for those countries with 'IX' in column 6 'Specific conditions' of Part 1 of Annex I to Regulation (EU) No 206/2010.
- (7) To be carried out according to the standards laid down in Annex III to Decision 2008/185/EC. In the case of pigs aged over 4 months, the test used shall be the whole virus ELISA.
- (8) Further requirements requested by Finland in respect of transmissible gastro-enteritis.

Official veterinarian	
Name (in capital letters):	Qualification and title:
Date:	Signature:
Stamp:	

	Model POR-Y							
		UNTRY		T		Veterinary cer	tificate to EU	
	I.1.	Consignor		I.2. Certific	ate reference num	ber I.2.a.		
		Name		I.3. Central Competent Authority				
		Address		I4 Local C	ompetent Authorit	·v		
		Tel. No				· y		
ent	1.5.	Consignee		I.6.				
n us		Name						
nsić		Address						
o p		Postal code						
tche		Tel. No	Г					
Part I: Details of dispatched consignment	1.7.	Country ISO code	I.8. Region Code of origin	I.9. Country destina		I.10. Region of destination	Code	
ils o	1.11	. Place of origin		I.12.				
l: Deta		Name Address	Approval number					
Part		Name Address	Approval number					
		Name Address	Approval number					
	I.13	. Place of loading Address	Approval number	I.14. Date of departure time of departure				
	I.15	. Means of transport Aeroplane Sh	nip	I.16. Entry BIP in EU				
		Road vehicle Oth Identification: Documentary references:	er 🗌	1.17.				
	I.18	. Description of commodity		I.19. Commodity code (HS code) 01.03				
					1.2	0. Quantity		
	I.21			I.22. Number of packages			es	
	1.23	3. Identification of container/s	eal number	1.24.				
	1.25	5. Commodities certified for: Slaughter						
	1.26	S.		I.27. For imp	ort or admission ir	nto EU		
	1.28	3. Identification of the commo	dities	1				
		Species (Scientific name)	Identification system	Identification number	ı	Age	Sex	

COUNTRY Model POR-Y II. Health information II.a. Certificate reference number II.b. II.1. **Public Health Attestation** I, the undersigned official veterinarian, hereby certify, that the animals described in this certificate: come from holdings which have been free from any official prohibition on health grounds, for the last 42 days in the case of brucellosis, for the last 30 days in the case of anthrax and for the past six months in the case of rabies and, Part II: Certification the animals have not been in contact with animals from holdings which did not satisfy these conditions; II.1.2 have not received: any stilbene or thyrostatic substances, oestrogenic, androgenic, gestagenic or β - agonist substances for purposes other than therapeutic or zootechnic treatment (as defined in Directive 96/22/EC). 11.2. **Animal Health attestation** I, the undersigned official veterinarian, hereby certify, that the animals described above meet the following requirements: they come from the territory with code:(1) which, at the date of issuing this certificate: (2) either [(a) has been free for 24 months from foot-and-mouth disease, for 12 months from rinderpest, African swine fever, classical swine fever, swine vesicular disease and vesicular exanthema, and for 6 months from vesicular stomatitis, and] (2) or [(a) (i) has been free [for 24 months from foot-and-mouth disease] (2), for 12 months from rinderpest, African swine fever, vesicular exanthema, [classical swine fever] (2) and [swine vesicular disease] (2), and for 6 months from vesicular stomatitis, and cases/outbreaks from that date, and authorised to export these animals by Commission (b) where during the last 12 months, no vaccination against these diseases has been carried out and imports of domestic cloven-hoofed animals vaccinated against these diseases are not they have remained in the territory described under point II.2.1 since birth, or for at least the last three months before 11.2.2 dispatch to the Union and without contact with imported cloven-hoofed animals for the last 30 days; they have remained in the holding(s) described under box reference I.11 since birth, or for at least 40 days prior to dispatch, and, during this period, in the holding(s) and in an area with a $10 \, \text{km}$ radius around the holding(s) of origin, 11.2.3 there has been no case/outbreak of the diseases referred to in point II.2.1; II.2.4 they are not animals to be killed under a national programme for the eradication of diseases, nor have they been vaccinated against the diseases referred to in point II.2.1; II.2.5 they are/were (2) dispatched from their holding(s) of origin, without passing through any market,

(2) either [directly to the Union,]

(2) or [to the officially authorised assembly centre described under box reference I.13 situated within the territory described under point II.2.1,]

and, until dispatched to the Union:

- (a) they did not come in contact with other cloven-hoofed animals not complying with the health requirements as described in this certificate, and
- (b) they were not at any place where, or around which within a 10 km radius, during the previous 40 days there has been a case/outbreak of any of the diseases referred to in point II.2.1;

COUNTRY Model POR-Y

II. Health information II.a. Certificate reference number II.b.

- II.2.6 any transport vehicles or containers in which they were loaded were cleaned and disinfected before loading with an officially authorised disinfectant;
- II.2.7 they were examined by an official veterinarian within 24 hours of loading and showed no clinical sign of disease;

II.3. Animal transport attestation

I, the undersigned official veterinarian, hereby certify, that the animals described above have been treated before and at the time of loading in accordance with the relevant provisions of Regulation (EC) No 1/2005, in particular as regards watering and feeding, and they are fit for the intended transport.

(2) (4) [II.4. Specific requirements

- II.4.1 Aujeszky's disease is notifiable in the country referred to in box reference I.7;
- II.4.2 according to official information, no clinical, pathological or serological evidence of Aujeszky's disease has been recorded in the holding(s) of origin referred to in box reference I.11, for the last 3 months;
- II.4.3 the animals referred to in box reference I.28:
 - (a) have remained in the holding(s) of origin referred to in box reference I.11 since birth or for the last 60 days prior to dispatch for exportation, and
 - (b) have not been vaccinated against Aujeszky's disease.]

Notes

This certificate is meant for live domestic porcine animals (Sus scrofa) intended for immediate slaughter after importation.

After importation the animals must be conveyed without delay to the slaughterhouse of destination to be slaughtered within five working days.

Part I:

- Box reference I.8: Provide the code of territory as appearing in Part 1 of Annex I to Regulation (EU) No 206/2010.
- Box reference I.13: The assembly centre, if any, must fulfil the conditions for its approval, as laid down in Part 5 of Annex I to Regulation (EU) No 206/2010.
- Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor must inform the BIP of entry into the Union.
- Box reference I.23: For containers or boxes, the container number and the seal number (if applicable) should be included.
- Box reference I.28: Identification system: The animals must bear:
 - An individual number which permits tracing of their premises of origin. Specify the identification system (such as tag, tattoos, brand, chip, transponder) and the anatomic place used in the animal.
 - An ear tag that includes the ISO code of the exporting country. The individual number must permit tracing of their premises of origin.
- Box reference I.28: Age: months.
- Box reference I.28: Sex (M = male, F = female, C = castrated).

CC	COUNTRY Model F							
II.	Health information	II.a. Certificate reference number	II.b.					
Pa	rt II:							
(1)	Code of the territory as it appears in Pa	rt 1 of Annex I to Regulation (EU) No 206/201	0.					
(2)	Keep as appropriate.							
(3)	(3) Date of loading. Imports of these animals shall not be allowed when the animals were loaded either prior to the date of authorisation for exportation to the Union of the third country, territory or part thereof referred to in boxes I.7 and I.8, or during a period where restrictive measures have been adopted by the Union against imports of these animals from this third country, territory or part thereof.							
(4)	When required by the EU Member Stat	e of destination, in accordance with Decision	2008/185/EC.					
Off	ficial veterinarian							
	Name (in capital letters):	Qualification	on and title:					
	Date:	Signature:						
	Stamp:							

	Model RUM							
		UNTRY		1			Veterinary ce	rtificate to EU
	I.1.	Consignor		I.2. Certific	ate reference	number	1.2.a.	
		Name		I.3. Central	Competent A	uthority		
		Address		L4. Local C	Competent Aut	thority		
		Tel. No						
ent	I.5.	Consignee		I.6.				
gnm		Name						
onsi		Address						
oo pe		Postal code						
tche		Tel. No						
Part I: Details of dispatched consignment	1.7.	Country ISC code	0	I.9. Country destina		SO ode	I.10. Region of destination	Code
ilso	1.11.	. Place of origin		I.12.				
: I: Deta		Name Address	Approval number					
Pari		Name Address	Approval number					
		Name Address	Approval number					
	I.13	. Place of loading Address	I.14. Date of departure time of departure					
	I.15	. Means of transport Aeroplane	I.16. Entry BIP in EU					
		Road vehicle	Other s:	I.17. No(s) of CITES				
	I.18	. Description of commodi	ty	I.19. Commodity code (HS code)				
				I.20. Quantity				
	I.21			I.22. Number of packages		es		
	1.23	. Identification of contain	er/seal number					
	1.25	i. Commodities certified fo	or:]	ı 🗆		Slau	ghter	
	1.26.			I.27. For imp	ort or admissi	ion into E	≣U	
	1.28	I. Identification of the com	modities					
		Species (Scientific name)	Identification system	Identification number	1	Ag	е	Sex

	COUN	TRY				Model RUI
	II.	Health	information		II.a. Certificate reference number	II.b.
	II.1.	Public	: Health Attest	ation		
I, the undersigned official veterinarian, hereby certify, that the animals described in this certificate:						cribed in this certificate:
II.1.1 come from a holding which has been free from any official prohibition on health grounds, for the last acase of brucellosis and tuberculosis, for the last 30 days in the case of anthrax, for the last six months rabies, and, have not been in contact with animals from holdings which did not satisfy these conditions. II.1.2 have not received: — any stilbene or thyrostatic substances, — pestrogenic androgenic gestagenic or 8- agonist substances for purposes other than therapeutic					e of anthrax, for the last six months in the case of	
ertific		II.1.2	have not rece	ved:		
± ∷C			any stilbe	ne or thyro	ostatic substances,	
Par					genic, gestagenic or β- agonist substances f ed in Directive 96/22/EC).	for purposes other than therapeutic or zootechnic
	II.2.	Anima	ıl Health Attes	ation		
		I, the ι	undersigned off	cial veterir	narian, hereby certify, that the animals desc	cribed above meet the following requirements:
II.2.1 they come from the territory with code:(1) which, at the date of issuing this certification.					which, at the date of issuing this certificate:	
 (a) has been free for 24 months from foot-and-mouth disease, for 12 months from rinderpest, blindered, contagious bovine pleuropneumonia, lumpy skin disease, peste des petits ruminants, pox, contagious caprine pleuropneumonia and epizootic haemorrhagic disease and for 6 mostomatitis, and (b) where during the last 12 months, no vaccination against these diseases has been carried cloven-hoofed animals vaccinated against these diseases are not permitted; 				, peste des petits ruminants, sheep pox and goat		
		11.2.2	they have ren	ained		
			(³) either	dispatc		birth, or for at least the last six months before oven-hoofed animals imported into this territory
			or	listed in condition country they ha	n Annex I, Part 7 to Regulation (EU) No 206 ons specified for each species in Annex I, Pa during a period of less than six months pr	e entry, if they are animals of the relevant species 1/2010 and they were imported directly under the eart 7 to Regulation (EU) No 206/2010 from a third rior to embarkation to the Union and in any case of the same health status after being released in a Union (2)]
		II.2.3	they have ren boxes referen			in the holding/establishment (3) described under
					n in an area of radius of 150 km, there has be ase during the previous 60 days, and	en no case/outbreak of bluetongue and epizootic
			` '		n in an area of 10 km radius, there has beer ing the previous 40 days;	n no case/outbreak of the other diseases referred
		II.2.4			be killed under a national programme for of the diseases referred to in point II.2.1, and	the eradication of diseases, nor have they been nd they:
			(³) (⁴) either	[come f	from a herd which is recognised as officially	tuberculosis free, and]
			(³) (⁵) or	[have b		ulin test within the past 30 days with negative

COUNTRY Model RUM

II.	Health	information	II.a. Certificate reference number	II.b.	
		they have not been vacc	inated against brucellosis and they:		
		(3) (4) either [come fr	om a herd which is recognised as officially bru	cellosis free;]	
		.,.,	een subjected to a serum agglutination test whic tination per ml, within the past 30 days;]	ch showed a brucella count of less than 30 IU	
		(3) or [are cas	trated males of any age;]		
	II.2.5	according to my knowled	ge and to the written declaration made by the owner, the animals:		
			holdings/establishments (3), and have not be ich the following diseases have been clinically		
		 contagious agalactia of sheep or goats (Mycoplasma agalactiae, Mycoplasma capricolum, Mycomycoides var. mycoides 'large colony'), within the last six months, 			
		(ii) paratuberculosis	and caseous lymphadenitis, within the last 12	months,	
		(iii) pulmonary aden	omatosis, within the last three years, and		
		(iv) Maedi/Visna or	caprine viral arthritis/encephalitis,		
		(³) either	[within the last three years,]		
			[within the last 12 months, and all the infected animals subsequently reacted negatively to apart,]		
		(b) are included in an of	ficial system for notification of these diseases,	and	
		(c) have been free from export;	clinical or other evidence of tuberculosis and	d brucellosis during the three years prior to	
	(³) (6) [II.2.6	(6) [II.2.6 the animals have reacted negatively to a serological test for the detection of antibody for bluetongue and epizootic haemorrhagic-disease, carried out on two occasions on samples of blood taken at the beginning of the isolation quarantine period and at least 28 days later on			
	II.2.7	they are dispatched fror Union and, until dispatch	n the holding/establishment described under lade to the Union:	boxes reference I.11 and I.13 directly to the	
		(a) they did not come ir described in this cer	contact with other cloven-hoofed animals not tificate, and	complying with the health requirements as	
			place where, or around which within a 10 km rak of any of the diseases referred to in point II.2		
	II.2.8	any transport vehicles or officially authorised disir	containers in which they were loaded were cle fectant;	aned and disinfected before loading with an	
	II.2.9	they were examined by a	an official veterinarian within 24 hours of loading	g and showed no clinical sign of disease;	
	II.2.10 they have been loaded for dispatch to the Union on			ed and disinfected before loading with an	
II.3.	. Anima	I transport attestation			
	I, the u	I, the undersigned official veterinarian, hereby certify, that the animals described above have been treated before and at the			

I, the undersigned official veterinarian, hereby certify, that the animals described above have been treated before and at the time of loading in accordance with the relevant provisions of Regulation (EC) No 1/2005, in particular as regards watering and feeding, and they are fit for the intended transport.

COUNTRY Model RUM

II. Health information II.a. Certificate reference number II.b.

(3) (8) [II.4. Specific requirements

- II.4.1 According to official information, no clinical or pathological evidence of infectious bovine rhinotracheitis (IBR) has been recorded in the holding/establishment (3) of origin referred to in boxes reference I.11 and I.13, for the last 12 months;
- II.4.2 the animals referred to in box reference I.28:
 - (a) have been isolated in accommodation approved by the competent authority for the last 30 days immediately prior to dispatch for export, and
 - (b) have been subjected to a serological test for IBR on sera taken at least 21 days after entry into isolation, with negative results, and all animals in isolation have also given negative results to this test, and
 - (c) have not been vaccinated against IBR.:

Notes

This certificate is meant for live animals of the order Artiodactyla (excluding bovine animals (including *Bubalus* and *Bison* species and their cross-breeds), *Ovis aries*, *Capra hircus*, Suidae and Tayassuidae), and of the families Rhinocerotidae and Elephantidae. Use one certificate per species.

After importation the animals must be conveyed without delay to the holding of destination where they shall remain for a minimum period of 30 days before further movement outside the holding, except in the case of a dispatch to a slaughterhouse.

Part I:

- Box reference I.8: Provide the code of territory as appearing in Part 1 of Annex I to Regulation (EU) No 206/2010.
- Box reference I.13: The assembly centre, if any, must fulfil the conditions for its approval, as laid down in Part 5 of Annex I to Regulation (EU) No 206/2010.
- Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor must inform the BIP of entry into the Union.
- Box reference I.19: Use the appropriate HS code: 01.02, 01.04.10, 01.04.20 or 01.06.19.
- Box reference I.23: For containers or boxes, the container number and the seal number (if applicable) should be included.
- Box reference I.28: Identification system: Specify the identification system (tag, tattoos, brand, chip, transponder). The ear tag
 includes the ISO code of the exporting country. The individual number must permit tracing of their premises of origin.
- Box reference I.28: Age: months.
- Box reference I.28: Sex (M = male, F = female, C = castrated).
- Box reference I.28: Species: Select the species amongst those listed for the following families:

Antilocapridae: Antilocapra spp.;

Bovidae: Addax spp., Aepyceros spp., Alcelaphus spp., Ammodorcas spp., Ammotragus spp., Antidorcas spp., Antilope spp., Boselaphus spp., Budorcas spp., Capra spp. (excluding Capra hircus), Cephalophus spp., Connochaetes spp., Damaliscus spp. (including Beatragus), Dorcatragus spp., Gazella spp., Hemitragus spp., Hippotragus spp., Kobus spp., Litocranius spp., Madoqua spp., Naemorhedus spp. (including Nemorhaedus and Capricornis), Neotragus spp., Oreamnos spp., Oreatragus spp., Orys spp., Ourebia spp., Ovis ospp., Vis spp. (excluding Ovis aries), Pantholops spp., Pelea spp., Procapra spp., Pseudois spp., Pseudoryx spp., Raphicerus spp., Redunca spp., Rupicapra spp., Saiga spp., Sigmocros-Alecelaphus spp., Sylvicapra spp., Syncerus spp., Taurotragus spp., Tetracerus spp., Tragelaphus spp. (including Boocerus).

 ${\it Camelidae: Camelus \ spp., \ Lama \ spp., \ Vicugna \ spp.}$

Cervidae: Alces spp., Axis-Hyelaphus spp., Blastocerus spp., Capreolus spp., Cervus-Rucervus spp., Dama spp., Elaphurus spp., Hippocamelus spp., Hydropotes spp., Mazama spp., Megamuntiacus spp., Muntiacus spp., Odocoileus spp., Ozotoceros spp., Pudu spp., Rangifer spp.

Giraffidae: Giraffa spp., Okapia spp.

Hippopotamidae: Hexaprotodon-Choeropsis spp., Hippopotamus spp.,

Moschidae: Moschus spp.

Tragulidae: *Hyemoschus* spp., *Tragulus-Moschiola* spp.,

Rhinocerotidae: Ceratotherium spp., Dicerorhinus spp., Diceros spp., Rhinoceros spp.

Elephantidae: Elephas spp., Loxodonta spp., as appropriate.

CC	COUNTRY Model RUM					
II.	Health information	II.a. Certificate reference number	II.b.			
(1) (2) (3) (4) (5)	art II: (¹) Code of the territory as it appears in Part 1 of Annex I to Regulation (EU) No 206/2010. (²) In this case the health certificate has to be accompanied by the official document on quarantine and test conditions laid down in Part 2 of Annex I to Regulation (EU) No 206/2010 (model 'CAM').					
Off	Official veterinarian					
	Name (in capital letters):	Qualificat	ion and title:			
	Date:	Signature				
	Stamp:	3				
	•					

			Mode	el SUI				
		UNTRY	1				Veterinary cer	tificate to EU
	l.1.	Consignor		I.2. Certifica	ate reference r	number	I.2.a.	
		Name		I.3. Central	Competent Au	uthority		
		Address		I.4. Local C	ompetent Auth	hority		
		Tel. No						
ent	1.5.	Consignee		1.6.				
gnm		Name						
onsi		Address						
) pe		Postal code						
tche		Tel. No						
Part I: Details of dispatched consignment	1.7.	Country ISO I.8. Reg of origin code of o	gion Code rigin	I.9. Country destinat		SO I	l.10. Region of destination	Code
sils	1.11	Place of origin		I.12.				
: I: Deta		Name Approva	ıl number					
Part		Name Approva	ıl number			/		
		Name Approva	ıl number					
	I.13	. Place of loading Address Approva	ıl number	I.14. Date of	departure	tin	ne of departure	
	I.15	Means of transport Aeroplane Ship	Railway wagon	I.16. Entry BI	P in EU			
		Road vehicle Other Identification: Documentary references:		I.17. No(s) of (CITES			
	I.18	. Description of commodity			I.19. Commo	odity cod	le (HS code)	
				l		I.20. Qı	uantity	
	I.21					I.22. Nu	umber of package	es
	1.23	. Identification of container/seal number	r			1.24.		
	1.25	. Commodities certified for:						
		Breeding	Fattening			Slaug	ghter 🗌	
	1.26			I.27. For impo	ort or admissio	on into E	U	
	1.28	. Identification of the commodities	-					
			fication stem	Identification number		Age	•	Sex

Part II: Certification

II. Health information II.a. Certificate reference number II.b.

II.1. Public Health Attestation

I, the undersigned official veterinarian, hereby certify, that the animals described in this certificate:

- III.1.1 come from a holding which has been free from any official prohibition on health grounds, for the last 42 days in the case of brucellosis, for the last 30 days in the case of anthrax and for the past six months in the case of rabies and, the animals have not been in contact with animals from holdings which did not satisfy these conditions;
- II.1.2 have not received:
 - any stilbene or thyrostatic substances,
 - oestrogenic, androgenic, gestagenic or β agonist substances for purposes other than therapeutic or zootechnic treatment (as defined in Directive 96/22/EC).

|| 2 Animal Health attestation

I, the undersigned official veterinarian, hereby certify, that the animals described above meet the following requirements:

- II.2.1 they come from the territory with code:(1) which, at the date of issuing this certificate:
 - (a) has been free for 24 months from foot-and-mouth disease, for 12 months from rinderpest, African swine fever, classical swine fever, swine vesicular disease and vesicular exanthema, and for 6 months from vesicular stomatitis. and
 - (b) where during the last 12 months, no vaccination against these diseases has been carried out and imports of cloven-hoofed animals vaccinated against these diseases are not permitted;
- II.2.2 they have remained in the territory described under point II.2.1 since birth, or for at least the last six months before dispatch to the Union and without contact with cloven-hoofed animals imported into this territory less than six months ago:
- II.2.3 they have remained in the holding described under boxes reference I.11 and I.13 since birth, or for 40 days prior to dispatch, and, during this period, in the holding(s) and in an area with a 10 km radius around the holding(s) of origin, there has been no case/outbreak of the diseases referred to in point II.2.1;
- II.2.4 A they are not animals to be killed under a national programme for the eradication of diseases, nor they have been vaccinated against the diseases referred to in point II.2.1 and they have been subjected within the past 30 days to a buffered Brucella antigen test for porcine brucellosis with negative results;
- (²) (³) [II.2.4 B they have been subjected within the past 30 days to a test for swine vesicular disease antibodies and a test for classical swine fever antibodies with negative results in both cases]
- (²) (4) [II.2.4 C they have been subjected within the past 30 days to a buffered Brucella antigen test for porcine brucellosis with negative results]
 - II.2.5 they come from holdings which:
 - (a) are not restricted under a national control and eradication programme for brucellosis, porcine enteroviral encephalomyelitis (Teschen disease), and
 - (b) are included in an official system for notification of these diseases;
 - II.2.6 they are dispatched from the holding described under boxes reference I.11 and I.13 directly to the Union and, until dispatched to the Union:
 - (a) they did not come in contact with other cloven-hoofed animals not complying with the health requirements as described in this certificate, and
 - (b) they were not at any place where, or around which within a 10 km radius, during the previous 40 days there has been a case/outbreak of any of the diseases referred to in point II.2.1;

COUN	TRY			Model SU			
II.	Health	information	II.a. Certificate reference number	II.b.			
	II.2.7	1.2.7 any transport vehicles or containers in which they were loaded were cleaned and disinfected before loading with an officially authorised disinfectant;					
	II.2.8	II.2.8 they were examined by an official veterinarian within 24 hours of loading and showed no clinical sign of disease;					
	II.2.9	2.9 they have been loaded for dispatch to the Union on					
II.3.	Anima	al transport attestation					
	I, the undersigned official veterinarian, hereby certify, that the animals described above have been treated before and at the time of loading in accordance with the relevant provisions of Regulation (EC) No 1/2005, in particular as regards watering and feeding, and they are fit for the intended transport.						
(²) (⁶) [I	I.4. Speci i	fic requirements					
	II.4.1	Aujeszky's disease is no	tifiable in the country referred to in box referen	ce I.7;			
	II.4.2	4.2 According to official information, no clinical, pathological or serological evidence of Aujeszky's disease has been recorded for the last 12 months in the holding(s) of origin referred to in boxes reference I.11 and I.13, and in an area with a 5 km radius around the holding(s);					
	II.4.3	the animals referred to in	box reference I.28:				
	(a) prior to dispatch for exportation, have remained since birth in the holding of origin referred to in the reference I.11 and I.13 or they have remained in this holding for the last 3 months and in others of equivi- status since birth,						
			in accommodation approved by the competer export, without direct or indirect contact with ot				
			d to an ELISA test for the presence of gl antil vith negative results; and, all animals in isolation				
(d) have not been vaccinated against Aujeszky's disease and have not been in the herd of origin has not been vaccinated during the previous 12 months.							
(2)	(8) [II.4.4]	(further requirements and/or tests)			
Notes							

This certificate is meant for live non-domestic Suidae (*Babyrousa* spp., *Hylochoerus* spp., *Phacochoerus* spp., *Potamochoerus* spp., and *Sus* spp.), Tayassuidae (*Catagonus* spp., *Pecari* spp., *Tayassu* spp.) and Tapiridae (*Tapirus* spp.).

After importation the animals must be conveyed without delay to the holding of destination where they shall remain for a minimum period of 30 days before further movement outside the holding, except in the case of a dispatch to a slaughterhouse.

COUNTRY Model SUI

II.	Health information	II.a. Certificate reference number	II.b.

Part I:

- Box reference I.8: Provide the code of territory as appearing in Part 1 of Annex I to Regulation (EU) No 206/2010.
- Box reference I.13: The assembly centre, if any, must fulfil the conditions for its approval, as laid down in Part 5 of Annex I to Regulation (EU) No 206/2010.
- Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor must inform the BIP of entry into the Union.
- Box reference I.19: Use the appropriate HS code: 01.03 or 01.06.19.
- Box reference I.23: For containers or boxes, the container number and the seal number (if applicable) should be included.
- Box reference I.28: Identification system: The animals must bear:
 - An individual number which permits tracing of their premises of origin. Specify the identification system (such as tag, tattoos, brand, chip, transponder) and the anatomic place used in the animal.
 - An ear tag that includes the ISO code of the exporting country. The individual number must permit tracing of their premises of origin.
- Box reference I.28: Age: months.
- Box reference I.28: Sex (M = male, F = female, C = castrated).
- Box reference I.28: Species.

Part II:

- (1) Code of the territory as it appears in Part 1 of Annex I to Regulation (EU) No 206/2010.
- (2) Keep as appropriate.
- (3) Supplementary guarantees to be provided when required in column 5 'SG' of Part 1 of Annex I to Regulation (EU) No 206/2010, with the entry 'B'.
- (4) Supplementary guarantees to be provided when required in column 5 'SG' of Part 1 of Annex I to Regulation (EU) No 206/2010, with the entry 'C'.
- (5) Date of loading. Imports of these animals shall not be allowed when the animals were loaded either prior to the date of authorisation for exportation to the Union of the third country, territory or part thereof referred to in boxes I.7 and I.8, or during a period where restrictive measures have been adopted by the Union against imports of Suidae animals from this third country, territory or part thereof.
- (6) When required by the EU Member State of destination, in accordance with Decision 2008/185/EC.
- (7) To be carried out according to the standards laid down in Annex III to Decision 2008/185/EC. In the case of animals aged over 4 months, the test used shall be the whole virus ELISA.
- (8) Further requirements requested by Finland in respect of transmissible gastro-enteritis.

Official veterinarian					
Name (in capital letters):	Qualification and title:				
Date:	Signature:				
Stamp:					

Model CAM Specific animal health attestation for animals quarantined in St. Pierre and Miquelon prior to introduction into the Union

	COUNTRY		Veterinary certificate to EU				
	I.1. Consignor		I.2. Certificat	te reference numbe	r I.2.a.		
	Name		I.3. Central C	I.3. Central Competent Authority			
	Address				,		
	Tel. No		I.4. Local Co	mpetent Authority			
Ħ	I.5. Consignee		1.6.				
nme	Name						
nsig	Address						
00 0	Postal code						
chec	Tel. No						
Part I: Details of dispatched consignment	I.7. Country ISO of origin code	I.8. Region Code of origin	l.9. Country of destination		I.10. Region of destination	Code	
o sli	I.11. Place of origin	·	I.12.				
Deta	Name	Approval number					
1::	Address						
Pal	Name Address	Approval number					
		Americal access on					
	Name Address	Approval number					
	I.13. Place of loading		I.14. Date of d	I.14. Date of departure time of departure			
	Address	Approval number					
	I.15. Means of transport Aeroplane Shi	p 🗌 Railway wagon 🗌	I.16. Entry BIF	P in EU			
	_	er 🗌	I.17. No(s) of CI	I.17. No(s) of CITES			
	Identification: Documentary references:						
	I.18. Description of commodity			I.19. Commodity c	ode (HS code) 0	1.06.19	
	ino. Becomplian or commodity			i.io. commodity o	040 (110 0040)		
				1.20.	Quantity		
	I.21.			1.22.	Number of packages		
	LOO Identification of a sately safe			104			
	I.23. Identification of container/se	eal number		1.24.			
	I.25. Commodities certified for:						
	Breeding	Fatten	ng 🗌	Sla	ughter		
	1.26.		I.27. For impo	rt or admission into	EU [
	I.28. Identification of the commod	dities					
	Species (Scientific name)	Identification system	Identification number	А	ge s	Sex	

COUNTRY Model CAM

Ĥ. Health information II.a. Certificate reference number II.b. II.1. Quarantine conditions attestation I, the undersigned official veterinarian, hereby certify, that the animals described in the animal health certificate (1) number Part 7 of Annex I to Regulation (EU) No 206/2010 for a period of: days before being released for exportation to the Part II: Certification Union and during this period they have been subject to the following tests (3), carried out in an approved laboratory within the Union, with a negative result (4): II.1.1. Brucellosis: (a) B. abortus: Serum Agglutination Test (SAT) and Rose Bengal Test (RBT) within two days after arrival and after at least 42 days (b) B. ovis: Complement Fixation Test (CFT) within two days after arrival and after at least 42 days (c) B. melitensis: SAT and RBT within two days after arrival and after at least 42 days II.1.2. Bluetongue and Epizootic haemorrhagic disease (5) either [two tests using Bluetongue competitive Elisa test within two days after arrival and after at least 21 days] [they have been quarantined for more than 60 days and during this period the quarantine station (5) or remained free of Bluetongue vectors (Culicoides), and no evidence of clinical disease has been detected]. II.1.3. Tuberculosis Two intradermal tuberculin test according to annex B to Directive 64/432/EC using bovine and avian tuberculin performed within two days after arrival and after at least 42 days from the first test II.1.4. Foot-and-mouth disease: ELISA test for the detection of antibodies and a virus neutralizaton test within two days after arrival and after at least 42 days II.1.5. Rinderpest: competitive ELISA test within two days after arrival and after at least 42 days II.1.6. Vesicular stomatitis: ELISA or virus- neutralisation test within two days after arrival and after at least 42 days II.1.7. Rift valley fever: an ELISA test or a virus neutralisation test within two days after arrival and after at least 42 days II.1.8. Lumpy skin disease: ELISA or virus neutralisation test within two days after arrival and after at least 42 days II.1.9. Crimean Congo haemorrhagic fever: ELISA or virus neutralisation test within two days after arrival and after at least 42 days II.1.10. Surra: blood microscopy within two days after arrival and after at least 42 days II.1.11. Malignant catarrhal fever: immunofluorescence test within two days after arrival and after at least 42 days II.2. Supplementary guarantees Bovine leukosis: AGID test or ELISA within two days after arrival and after at least 42 days (When required by the EU

Member State of destination) (5)

COUNTRY Model CAM

Health	information		II.a. Certificate reference	number	II.b.	
Treatments						
They h	They have been subjected to:					
II.3.1.	an internal and external antiparasitic treatment during the quarantine period			riod		
II.3.2.						
	(5) either	[a treatm	ent with streptomycin 25mg	ı/kg]		
	(5) or [an antibiotic treatment effective against Leptospira spp. (specifymg/kg		p. (specify			
(⁵) [II.3.3.					mm/yyyy) using vaccine	
	Treatm They h II.3.1.	They have been subject II.3.1. an internal and II.3.2. (5) either (5) or	Treatments They have been subjected to: II.3.1. an internal and external at II.3.2. (5) either [a treatm (5) or [an antib mg/kg (5) [II.3.3. a vaccination against rabi	Treatments They have been subjected to: II.3.1. an internal and external antiparasitic treatment during II.3.2. (5) either [a treatment with streptomycin 25mg (5) or [an antibiotic treatment effective agmg/kg	Treatments They have been subjected to: II.3.1. an internal and external antiparasitic treatment during the quarantine per II.3.2. (5) either [a treatment with streptomycin 25mg/kg] (6) or [an antibiotic treatment effective against Leptospira sping/kg	Treatments They have been subjected to: II.3.1. an internal and external antiparasitic treatment during the quarantine period II.3.2. (5) either [a treatment with streptomycin 25mg/kg] (5) or [an antibiotic treatment effective against Leptospira spp. (specifymg/kg

Notes

This certificate is meant for live animals of the family Camelidae.

Part I:

- Box reference I.8: Provide the code of territory as appearing in Part 1 of Annex I to Regulation (EU) No 206/2010.
- Box reference I.13: The assembly centre, if any, must fulfil the conditions for its approval, as laid down in Part 5 of Annex I to Regulation (EU) No 206/2010.
- Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor must inform the BIP of entry into the Union.
- Box reference I.23: For containers or boxes, the container number and the seal number (if applicable) should be included.
- Box reference I.28: Identification system: The animals must bear:
 - An individual number which permits tracing of their premises of origin. Specify the identification system (such as tag, tattoos, brand, chip, transponder) and the anatomic place used in the animal.
 - An ear tag that includes the ISO code of the exporting country. The individual number must permit tracing of their premises of origin.
- Box reference I.28: Age: months.
- Box reference I.28: Sex (M = male, F = female, C = castrated).
- Box reference I.28: Species: Select amongst 'Camelus spp.', 'Lama spp.', 'Vicugna spp.' as appropriate.

Part II:

- (¹) Animal health certificate for non domestic animals other than Suidae, consigned to the Union (model 'RUM') as laid down in Part 2 of Annex I to Regulation (EU) No 206/2010.
- (2) Date in which the last animal in a group entered the quarantine facility.
- (3) Tests performed in accordance with the methods described in Chapter 2 of Part 7 of Annex I to Regulation (EU) No 206/2010.
- (4) Results of the tests performed must be attached in original to this health attestation.
- (5) Keep as appropriate.

NB:Sampling and testing procedures must be grouped as much as possible while respecting the minimum time intervals to avoid excessive handling and manipulation of the animals.

COUNTR	RY		Model CAM
II.	Health information	II.a. Certificate reference number	II.b.
Official ve	eterinarian		
	Name (in capital letters):	Qualification	and title:
	Date:	Signature:	
	Stamp		

PART 3

Addendum for transport of animals by sea

(To be completed and attached to the veterinary certificate when transport to the Union frontier includes, even for part of the journey, transportation by ship.)

Declaration by the master of the ship				
I, the undersigned, master of ship (name), declare that the animals referred to in the attached veterinary certificate No				
Done at	on			
(Port of arrival)	(Date of arrival)			
	(signature of master)			
(stamp)				
	(name in capital letters and title)			

PART 4

Addendum for transport of animals by air

(To be completed and attached to the veterinary certificate when transport to the Union frontier includes, even for part of the journey, transportation by air.)

Declaration by the captain of the aircraft	
I, the undersigned, captain of the aircraft (name container and the area around the crate or container attached veterinary certificate No has departure.	containing the animals referred to in the
Done at on	
(Airport of departure)	(Date of departure)
	(signature of captain)
(stamp)	
	(name in capital letters and title)

PART 5

Conditions for the approval of assembly centres (referred to in Article 4)

In order to be approved, assembly centres must meet the following requirements:

- I. They must be supervised by an official veterinarian.
- II. They must each be situated at the centre of an area of at least 20 km in diameter in which, according to official findings, there has been no case of foot-and-mouth disease for at least a period of 30 days prior to their use as approved assembly centres.

▼C1

- III. They must, before each use as approved assembly centres, be cleansed and disinfected with a disinfectant officially authorised in the exporting country as effective for the control of foot-and-mouth disease.
- IV. They must have, taking into account their animal capacity:
 - (a) a facility dedicated exclusively for use as an assembly centre;
 - (b) appropriate facilities, that are easy to clean and disinfect, for loading, unloading and adequate housing of a suitable standard for the animals, for watering and feeding them, and for giving them any necessary treatment;
 - (c) appropriate facilities for inspection and isolation;
 - (d) appropriate equipment for cleaning and disinfecting rooms and trucks;
 - (e) an appropriate storage area for fodder, litter and manure;
 - (f) an appropriate system for collecting and disposal of waste water;
 - (g) an office for the official veterinarian.
- V. When operating, they must have sufficient veterinarians to carry out all duties set out in Part 5;
- VI. They must only admit animals that are individually identified so as to guarantee traceability. To this end, when animals are admitted the owner or the person in charge of the centre must ensure that the animals are properly identified and accompanied by health documents or certificates for the species and categories involved.

In addition, the owner or the person in charge of the assembly centre must record on a register or in a data base, and retain for at least three years the name of the owner, the origin of the animals, the dates of entry and exit, the identification number of the animals or registration number of the herd of origin and the holding of destination, and, the registration number of the carrier and the registration number of the lorry delivering or collecting animals from that assembly centre.

- VII. All animals passing through the assembly centre must fulfil the health conditions established for the introduction of the relevant category of animal into the Union.
- VIII. Animals to be introduced into the Union which pass through an assembly centre must, within six days of arrival at the assembly centre, be loaded and dispatched directly to the border of the exporting country:
 - (a) without coming into contact with cloven-hoofed animals other than animals which fulfil the health conditions established for the introduction of the relevant category of animal into the Union;
 - (b) segregated into consignments so that no consignment contains both animals for breeding or production and animals for immediate slaughter;
 - (c) in transport vehicles or containers which have first been cleansed and disinfected with a disinfectant officially authorised in the exporting country as effective for the control of foot-and-mouth disease and which are so constructed that faeces, urine, litter or fodder cannot flow or fall out during transportation.

▼C1

- IX. Where the conditions for the export of animals to the Union require that a test is carried out within a specified period before loading, that period must include any period of assembly, up to six days, from the date of arrival of the animals at the approved assembly centre.
- X. The exporting third country must designate the centres which are approved for animals for breeding and production and those centres which are approved for animals for slaughter and must notify the Commission and the competent central authorities of the Member States of the names and addresses of such premises. That information must be updated regularly.
- XI. The exporting third country shall determine the procedure for official supervision of approved assembly centres and shall ensure that such supervision is carried out.
- XII. The approved assembly centres must be regularly inspected by the competent authority of the third country in order to check that the requirements for approval set out in points I to XI continue to be fulfilled.

If those inspections show that those conditions are no longer complied with, the approval of the centre must be suspended. The approval may be restored only when the competent authority of the third country is satisfied that the centre fully complies with the conditions set out in points I to XI.

PART 6

Protocols for the standardisation of materials and testing procedures

(referred to in Article 5)

Tuberculosis (TBL)

The single intradermal tuberculin test using bovine tuberculin shall be carried out according to Annex B to Directive 64/432/EEC. In the case of Suidae animals, the single intradermal tuberculin test using avian tuberculin shall be carried out according to Annex B to 64/432/EEC, except that the site of injection shall be the loose skin at the base of the ear.

▼<u>M2</u>

Brucellosis (Brucella abortus) (BRL)

The serum agglutination test, complement fixation test, buffered brucella antigen test, enzyme-linked immunosorbent assays (ELISA) and fluorescence polarisation assay (FPA) shall be carried out according to Annex C to Directive 64/432/EEC.

▼<u>C1</u>

Brucellosis (Brucella melitensis) (BRL)

Tests shall be carried out according to Annex C to Directive 91/68/EEC.

Enzootic Bovine Leukosis (EBL)

The agar gel immuno-diffusion test and the enzyme linked immuno-absorbent assay test (ELISA) shall be carried out according to paragraphs A and C of Chapter II of Annex D to Directive 64/432/EEC.

Bluetongue (BTG)

A. The blocking or competitive ELISA test shall be carried out according to the following protocol:

The competitive ELISA using monoclonal antibody 3-17-A3 is capable of identifying antibodies to all known serotypes of bluetongue virus (BTV).

The principle of the test is the interruption of the reaction between BTV antigen and a group-specific monoclonal antibody (3-17-A3) by the addition of test serum. Antibodies to BTV present in the test serum block the reactivity of the monoclonal antibody (Mab) and result in a reduction in the expected colour development after the addition of enzyme labelled anti-mouse antibody, and chromogen/ substrate. Sera can be tested at a single dilution of 1:5 (spot test - Appendix 1) or may be titrated (serum titration - Appendix 2) to give dilution end-point. Inhibition values higher than 50 % may be regarded as positive.

Material and Reagents:

- 1. Appropriate ELISA microtitre plates.
- 2. Antigen: supplied as a cell extracted concentrate, prepared as described below, and stored at either - 20 °C or - 70 °C.
- 3. Blocking buffer: phosphate buffered saline (PBS) containing 0,3 % BTV negative adult bovine serum, 0,1 % (v/v) Tween-20 (supplied as polyoxyethylene sorbiton monolaurate syrup) in PBS.
- 4. Monoclonal antibody: 3-17-A3 (supplied as hybridoma tissue-culture supernatant) directed against the group-specific polypeptide VP7, stored at - 20 °C or freeze-dried and diluted 1/100 with blocking buffer before use.
- 5. Conjugate: rabbit anti-mouse globulin (adsorbed and eluted) conjugated to horseradish peroxidase and kept in the dark at 4 °C.
- 6. Chromogen and substrate: Orthophenylene diamine (OPD-chromogen) at a final concentration of 0,4 mg/ml in sterile distilled water. Hydrogen peroxide (30 %w/v-substrate) 0,05 % v/v added immediately before use (5μl H₂ O₂ per 10 ml OPD). (Handle OPD with care - wear rubber gloves - suspected mutagen).
- 7. 1 Molar sulphuric acid: 26,6 ml of acid added to 473,4 ml of distilled water. (Remember Acid must be added to water, never water to acid.)
- 8. Orbital shaker.
- 9. ELISA plate reader (the test may be read visually).

Test format

Cc: conjugate control (no serum/ no monoclonal antibody); C++: strong positive serum control; C+: weak positive serum control; C-: negative serum control; Cm: monoclonal antibody control (no serum).

APPENDIX 1:

Spot dilution (1:5) format (40 sera/plate)

	Con	trols	Test Sera									
	1	2	3	4	5	6	7	8	9	10	11	12
A	Сс	C-	1	2	3	4	5	6	7	8	9	10
В	Сс	C-	1	2	3	4	5	6	7	8	9	10

	Con	trols	Test Sera									
	1	2	3	4	5	6	7	8	9	10	11	12
С	C++	C++										
D	C++	C++										
Е	C+	C+										
F	C+	C+										
G	Cm	Cm										40
Н	Cm	Cm										40

APPENDIX 2:

Serum titration format (10 sera/plate)

	Con	trols	Test Sera									
	1	2	3	4	5	6	7	8	9	10	11	12
A	Cc	C-	1:5									1:5
В	Сс	C-	1:10									1:10
С	C++	C++	1:20									1:20
D	C++	C++	1:40									1:40
Е	C+	C+	1:80									1:80
F	C+	C+	1:160									1:160
G	Cm	Cm	1:320									1:320
Н	Cm	Cm	1:640									1:640

Test protocol:

Conjugate control (Cc): Wells 1A and 1B are a blank control consisting of

BTV antigen and conjugate. This may be used to

blank the ELISA reader.

Mab control (Cm): Columns 1 and 2, rows G and H are the mono-

clonal antibody control and contain BTV antigen, monoclonal antibody and conjugate. These wells represent maximum colour. The mean of the optical density readings from this control

represents the 0 % inhibition value.

Positive control (C++, C+): Columns 1 and 2, rows C-D-E-F. These wells contain BTV antigen, BTV strong and weak positive antiserum respectively, Mab and

conjugate.

Negative control (C-): Wells 2A and 2B are the negative controls, which

contain BTV antigen, BTV negative antiserum,

Mab and conjugate.

Test sera: For large-scale serological surveys and rapid

screening, sera may be tested at a single dilution of 1:5 (Appendix 1). Alternatively, 10 sera may be tested over a dilution range from 1:5 to 1:640 (Appendix 2). This will give some indication of

the titre of antibody in the test sera.

Procedure:

1. Dilute BTV antigen to pre-titrated concentration in PBS, sonicate briefly to disperse aggregated virus (if sonicator is not available, pipette vigorously) and add 50 μ l to all wells of the ELISA plate. Tap sides of plate to disperse antigen.

- Incubate at 37 °C for 60 minutes on an orbital shaker. Wash plates three times by flooding and emptying the wells with non-sterile PBS and blot dry on absorbent paper.
- Control wells: Add 100 μl of blocking buffer to Cc wells. Add 50 ul of positive and negative control sera, at a dilution of 1:5 (10 μ l sera + 40 μl blocking buffer), to respective wells C-, C+ and C++. Add 50μl blocking buffer to Mab control wells.

Spot titration method: Add a 1:5 dilution of each test serum in blocking buffer to duplicate wells of columns 3 to 12 (10 μ l sera + 40 μ l blocking buffer),

or

Serum titration method: Prepare a two-fold dilution series of each test sample (1:5 to 1:640) in blocking buffer across eight wells of single columns 3 to 12.

- 4. Immediately after the addition of the test sera, dilute Mab 1:100 in blocking buffer and add 50 μ l to all wells of the plate except for the blank control.
- Incubate at 37 °C for 60 minutes on an orbital shaker. Wash three times with PBS and blot dry.
- Dilute rabbit anti-mouse concentrate to 1/5 000 in blocking buffer and add 50
 μl to all wells of the plate.
- Incubate at 37 °C for 60 minutes on an orbital shaker. Wash three times with PBS and blot dry.
- 8. Thaw the O-Phenylenediamine dihydrochloride (OPD) and immediately before use add 5 μl of 30 % hydrogen peroxide to each 10 ml of OPD. Add 50 μl to all wells of the plate. Allow colour to develop for approximately 10 minutes and stop the reaction with 1 Molar sulphuric acid (50 μl per well). Colour should develop in the Mab control wells and in those wells containing sera with no antibody to BTV.

Examine and record the plates either visually or using a spectrophotometric reader.

Analysis of results:

Using the software package print out the optical density (OD) values, and the percentage inhibition (PI) for test and control sera based on the mean value recorded in the antigen control wells. The date expressed as OD and PI values are used to determine whether the test has performed within acceptable limits. The upper control limits (UCL) and lower control limits (LCL) for the Mab control (antigen plus Mab in the absence of test sera) are between OD values 0,4 and 1.4. Any plate that fails to conform to the above criteria must be rejected.

If a computer software package is not available print out the OD values using the ELISA printer. Calculate the mean OD value for the antigen control wells, which is equivalent to the 100 % value. Determine the 50 % OD value and manually calculate the positivity and negativity of each sample.

Percentage inhibition (PI) value = 100 – (OD of each test control/Mean OD of Cm) \times 100.

The duplicate negative control serum wells and the duplicate blank wells must record PI values between + 25 % and - 25 %, and between + 95 % and + 105 %, respectively. Failure to be within these limits does not invalidate the plate but does suggest that background colour is developing. The strong and weak positive control sera must record PI values between + 81 % and + 100 %, and between + 51 % and + 80 %, respectively.

The diagnostic threshold for test sera is 50 % (PI 50 % or OD 50 %). Samples recording PI values >50 % are recorded negative. Samples that record PI values above and below the threshold for the duplicate wells are considered doubtful; such samples may be re-tested in the spot test and/or titration. Positive samples may also be titrated to provide an indication of the degree of positivity.

Visual reading: Positive and negative samples are easily discernible by eye; weakly positive or strong negative samples may be more difficult to interpret by eye.

Preparation of BTV ELISA antigen:

- 1. Wash 40-60 roux of confluent BHK-21 cells three times with serum-free Eagle's medium and infect with bluetongue virus serotype 1 in serum-free Eagle's medium.
- 2. Incubate at 37 °C and examine daily for cytopathic effect (CPE).
- 3. When CPE are complete in 90 % to 100 % of the cell sheet of each roux, harvest the virus by shaking any still-attached cells from the glass.
- 4. Centrifuge at 2 000 to 3 000 rpm to pellet the cells.
- 5. Discard the supernatant and re-suspend the cells in approximately 30 ml of PBS containing 1 % 'Sarkosyl' and 2 ml phenylmethylsulphonyl fluoride (lysis buffer). This may cause the cells to form a gel and more lysis buffer may be added to reduce this effect. (NB: phenylmethylsulphonyl fluoride is harmful - handle with extreme caution.)
- Disrupt the cells for 60 seconds using an ultrasonic probe at an amplitude of 30 microns.

- 7. Centrifuge at 10 000 rpm for 10 minutes.
- 8. Store the supernatant at + 4 °C and re-suspend the remaining cell pellet in 10 to 20 ml of lysis buffer.
- Sonicate and clarify, storing the supernatant at each stage, a total of three times
- 10. Pool the supernatants and centrifuge at 24 000 rpm (100,000 g) for 120 minutes at + 4 °C over a 5 ml cushion of 40 % sucrose (w/v in PBS) using 30 ml Beckmann centrifuge tubes and an SW 28 rotor.
- 11. Discard the supernatant, drain the tubes thoroughly and re-suspend the pellet in PBS by sonication. Store the antigen in aliquots at -20 °C.

Titration of BTV ELISA antigen:

Bluetongue ELISA antigen is titrated by the indirect ELISA. Twofold dilutions of antigen are titrated against a constant dilution (1/100) monoclonal antibody 3-17-A3. The protocol is as follows:

- 1. Titrate a 1:20 dilution of BTV antigen in PBS across the microtitre plate in a twofold dilution series (50 μ l/well) using a multichannel pipette.
- 2. Incubate for one hour at 37 °C on an orbital shaker.
- 3. Wash plates three times with PBS.
- Add 50 µl of monoclonal antibody 3-17-A3 (diluted 1/100) to each well of the microtitre plate.
- 5. Incubate for one hour at 37 °C on an orbital shaker.
- 6. Wash plates three times with PBS.
- 7. Add 50 μ l of rabbit anti-mouse globulin conjugated to horseradish peroxidase, diluted to a pre-titrated optimal concentration, to each well of the microtitre plate.
- 8. Incubate for one hour at 37 °C on an orbital shaker.
- Add substrate and chromogen as described previously. Stop the reaction after 10 minutes by the addition of 1 Molar sulphuric acid (50 μl/well).

In the competitive assay, the monoclonal antibody must be in excess, therefore a dilution of antigen is chosen which falls on the titration curve (not on the plateau region) which gives approximately 0,8 OD after 10 minutes.

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B. The agar gel immuno-diffusion test shall be carried out according to the following protocol:

Antigen:

Precipitating antigen is prepared in any cell culture system that supports the rapid multiplication of a reference strain of bluetongue virus. BHK or Vero cells are recommended. Antigen is present in the supernatant fluid at the end of virus growth but requires 50 to 100-fold concentration to be effective. This may be achieved by any standard protein concentration procedure; virus in the antigen may be inactivated by the addition of 0.3~%~(v/v) beta-propiolactone.

Known positive control serum:

Using the international reference serum and antigen a national standard serum is produced, standardised for optimal proportion against the international reference serum, freeze-dried and used as the known control serum in each test.

Test serum

Procedure:

1 % agarose prepared in borate or sodium barbitol buffer, pH 8,5 to 9,0, is poured into a petri dish to a minimum depth of 3,0 mm. A test pattern of seven moisture-free wells, each 5,0 mm in diameter, is cut in the agar. The pattern consists of one centre well and six wells arranged round it in a circle of radius 3 cm. The central well is filled with the standard antigen. Peripheral wells 2, 4 and 6 are filled with known positive serum, wells 1, 3 and 5 are filled with test sera. The system is incubated for up to 72 hours at room temperature in a closed humid chamber.

Interpretation:

A test serum is positive if it forms a specific precipitin line with the antigen and forms a complete line of identity with the control serum. A test serum is negative if it does not form a specific line with the antigen and it does not bend the line of the control serum. Petri dishes must be examined against a dark background and using indirect illumination.

Epizootic haemorrhagic disease (EHD)

The agar gel immuno-diffusion test shall be carried out according to the following protocol:

Antigen:

Precipitating antigen is prepared in any cell culture system that supports the rapid multiplication of the appropriate serotype(s) of epizootic haemorrhagic disease virus. BHK or Vero cells are recommended. Antigen is present in the supernatant fluid at the end of virus growth but requires 50 to 100-fold concentration to be effective. This may be achieved by any standard protein concentration procedure; virus in the antigen may be inactivated by the addition of 0.3% (v/v) beta-propiolactone.

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Known positive control serum:

Using the international reference serum and antigen a national standard serum is produced, standardised for optimal proportion against the international reference serum, freeze-dried and used as the known control serum in each test.

Test serum

Procedure:

1 % agarose prepared in borate or sodium barbitol buffer, pH 8,5 to 9,0, is poured into a petri dish to a minimum depth of 3,0 mm. A test pattern of seven moisture-free wells, each 5,0 mm in diameter, is cut in the agar. The pattern consists of one centre well and six wells arranged round it in a circle of radius 3 cm. The central well is filled with the standard antigen. Peripheral wells 2, 4 and 6 are filled with known positive serum, wells 1, 3 and 5 are filled with test sera. The system is incubated for up to 72 hours at room temperature in a closed humid chamber.

Interpretation:

A test serum is positive if it forms a specific precipitin line with the antigen and forms a complete line of identity with the control serum. A test serum is negative if it does not form a specific line with the antigen and it does not bend the line of the control serum. Petri dishes must be examined against a dark background and using indirect illumination.

Infectious bovine rhinotracheitis (IBR) / infectious pustular vulvo-vaginitis (IPV)

A. The serum neutralisation test shall be carried out according to the following protocol:

Serum: All sera are heat-inactivated at 56 °C for 30 minutes

before use.

Procedure: The constant virus-varying serum neutralisation test on

microtitre plates employs MDBK or other susceptible cells. The Colorado, Oxford or any other reference strain of the virus is used at 100 TCID50 per 0,025 ml; inactivated undiluted serum samples are mixed with an equal volume (0,025 ml) of virus suspension. The virus/serum mixtures are incubated for 24 hours at 37 °C in the microtitre plates before the MDBK cells are added. Cells are used at a concentration which

forms a complete monolayer after 24 hours.

Controls: (i) virus infectivity assay, (ii) serum toxicity controls,

(iii) uninoculated cell culture controls, (iv) reference

antisera.

Interpretation: The results of the neutralisation test and the titre of the

virus used in the test are recorded after three to six days incubation at 37 °C. Serum titres are considered negative if there is no neutralisation at a dilution of

1/2 (undiluted serum).

B. Any other test recognised in the framework of Decision 2004/558/EC (1).

⁽¹⁾ OJ L 249, 23.7.2004, p. 20.

Foot-and-mouth disease (FMD)

A. Collecting oesophageal/pharyngeal samples and testing shall be carried out according to the following protocol:

Reagents:

Prior to sampling, transport medium is prepared. Two ml volumes are dispensed in as many containers as there are animals to be sampled. The containers used must withstand freezing over solid CO2 or liquid nitrogen. Samples are obtained by the use of a specially-designed sputum collector or 'probang'. To obtain a sample the probang cup is passed through the mouth, over the dorsum of the tongue and down into the upper part of the oesophagus. Attempts are made to scrape the surface epithelium of the upper oesophagus and pharynx by movements directed laterally and dorsally. The probang is then withdrawn, if possible after the animal has swallowed. The cup must be full and contain a mixture of mucus, saliva, oesophageal fluid and cellular debris. Care must be taken to ensure that each specimen contains some visible cellular material. Very rough handling which causes bleeding must be avoided. Samples from some animals may be heavily contaminated with ruminal contents. Such samples must be discarded and the mouth of the animal flushed with water, or preferably physiological saline, before repeat sampling.

Treatmentof samples::

Each sample collected in the probang cup is examined for quality and 2 ml added to an equal volume of transport medium in a container which can withstand freezing. The containers are tightly closed, sealed, disinfected and labelled. The samples are kept cool (+ 4 °C) and examined within three to four hours or placed over dry ice (- 69 °C) or liquid nitrogen and kept frozen until examined. Between animals the probang is disinfected and washed in three changes of clean water.

Testing for FMD virus::

Samples are inoculated into cultures of primary bovine thyroid cell cultures using at least three tubes per sample. Other susceptible cells such as primary bovine or porcine kidney cells can be used but it must be kept in mind that for some strains of FMD virus they are less sensitive. The tubes are incubated at 37 °C on a roller apparatus and examined daily for 48 hours for the presence of a cytopathic effect (CPE). If negative, cultures are blind passaged onto new cultures and re-examined for 48 hours. The specificity of any CPE must be confirmed.

Recommended transport media:

- 0,08M phosphate buffer pH 7,2 containing 0,01 % bovine serum albumin, 0,002 % phenol red and antibiotics.
- 2. Tissue culture medium (such as Eagle's MEM) containing 0,04 M Hepes buffer, 0,01 % bovine serum albumin and antibiotics, pH 7,2.
- Antibiotics (per ml final) must be added to the transport medium such as penicillin 1 000 IU, neomycin sulphate100 IU, polymyxin B sulphate50 IU, mycostatin100 IU.
- B. The virus neutralisation test shall be carried out according to the following protocol:

Reagents:

Stock FMDV antigen is prepared in cell cultures or on cattle tongues and stored at - 70 °C or less or at - 20 °C after the addition of 50 % glycerol. This is the stock antigen. FMDV is stable under these conditions and titres vary little over a period of months.

Procedure:

The test is carried out in flat-bottomed tissue culture grade microtitre plates using susceptible cells such as IB-RS-2, BHK-21 or calf kidney cells. Sera for the test are diluted 1/4 in serum-free cell culture medium with the addition of 100 IU/ml neomycin or other suitable antibiotics. Sera are inactivated at 56 °C for 30 minutes and 0.05 ml amounts are used to prepare a twofold series on microtitre plates using 0,05 ml diluting loops. Pre-titrated virus also diluted in serum-free culture medium and containing 100 TCID50/0.05 ml is then added to each well. Following incubation at 37 °C for one hour to allow neutralisation to take place, 0,05 ml of suspension cells containing 0,5 to 1.0×10^6 cells per 1 ml in cell culture medium containing serum free of FMD antibody is added to each well and the plates are sealed. Plates are incubated at 37 °C. Monolayers are normally confluent within 24 hours. CPE is usually sufficiently advanced at 48 hours for a microscopic reading of the test. At this time a final microscopic reading may be made or the plates may be fixed and stained for macroscopic reading, for instance using 10 % formol-saline and 0,05 % methylene blue.

Controls:

Controls in each test include homologous antiserum of known titre, a cell control, a serum toxicity control, a medium control, and a virus titration from which the actual amount of virus in the test is calculated.

Interpretation:

Wells with evidence of CPE are considered to be infected and neutralisation titres are expressed as the reciprocal of the final dilution of serum present in the serum/virus mixtures at the 50 % end point estimated according to the Spearman-Karber method. (Karber, G., 1931, Archiv fuer Experimentelle Pathologie und Pharmokologie, 162, 480.). Tests are considered to be valid when the actual amount of virus used per well in the test is between 101,5 and 102,5 TCID50 and when the titre of the reference serum is within twofold of its expected titre, estimated from the mode of previous titrations. When the controls are outside these limits the tests are repeated. An end point titre of 1/11 or less is taken as negative.

C. The detection and quantification of antibody by ELISA shall be carried out according to the following protocol:

Reagents:

Rabbit antisera to 146S antigen of seven types of foot-and-mouth disease virus (FMDV) used at a predetermined optimum concentration carbonate/bicarbonate buffer, pH 9,6. Antigens are prepared from selected strains of virus grown on monolayers of BHK-21 cells. The unpurified supernatants are used and pretitrated according to the protocol but without serum, to give a dilution which after the addition of an equal volume of PBST (phosphate buffered saline containing 0,05 % Tween-20 and phenol red indicator) would give an optical density reading of between 1,2 and 1,5. The viruses can be used inactivated. PBST is used as a diluent. Guinea-pig antisera are prepared by inoculating guinea pigs with 146S antigen of each serotype. A predetermined optimum concentration is prepared in PBST containing 10 % normal bovine serum and 5 % normal rabbit serum. Rabbit anti-guinea-pig immunoglobulin conjugated to horseradish peroxidase is used at a predetermined optimum concentration in PBST containing 10 % normal bovine serum and 5 % normal rabbit serum. Test sera are diluted in PBST.

Procedure:

- 1. ELISA plates are coated with 50 μl of rabbit antiviral sera overnight in a humidity chamber at room temperature.
- 2. Fifty microlitres of a duplicate, twofold series of each test serum starting at 1/4 are prepared in U-bottomed multiwell plates (carrier plates). Fifty microlitres of a constant dose of antigen are added to each well and the mixtures are left overnight at 4 °C. The addition of the antigen reduces the starting serum dilution to 1/8.
- 3. The ELISA plates are washed five times with PBST.
- 4. Fifty microlitres of serum/antigen mixtures are then transferred from the carrier plates to the rabbit-serum-coated ELISA plates and incubated at 37 °C for one hour on a rotary shaker.

- After washing, 50 µl of guinea-pig antiserum to the antigen used in point 4 is added to each well. The plates are incubated at 37 °C for one hour a rotary shaker.
- 6. The plates are washed and 50 μ l of rabbit anti-guinea-pig immunoglobulin conjugated to horseradish peroxidase is added to each well. The plates are incubated at 37 °C for one hour on a rotary shaker.
- 7. The plates are washed and 50 μ l of orthophenylene diamine containing 0,05 % H_2O_2 (30 %) w/v is added to each well.
- 8. The reaction is stopped after 15 minutes with 1,25M H₂SO₄.

The plates are read spectrophotometrically at 492 nm on an ELISA reader linked to a microcomputer.

Controls: For each antigen used 40 wells contain no serum

but contain antigen diluted in PBST. A duplicated twofold dilution series of homologous bovine reference antiserum. A duplicate twofold dilution

series of negative bovine serum.

Interpretation: Antibody titres are expressed as the final dilution

of tests serum giving 50 % of the mean OD value recorded in the virus control wells where test serum is absent. Titres in excess of 1/40 are

considered positive.

References: Hamblin C, Barnett ITR and Hedger RS (1986) 'A

new enzyme-linked immunosorbent assay (ELISA) for the detection of antibodies against foot-and-mouth disease virus. I. Development and method of ELISA.' Journal of Immunological Methods, 93,

115 to 121.11.

Aujeszky's disease (AJD)

A. The serum neutralisation test shall be carried out according to the following protocol:

Serum: All sera are heat-inactivated at 56 °C for 30

minutes before use.

Procedure: The constant virus-varying serum neutralisation

test on microtitre plates employs Vero or other sensitive cell systems. Aujeszky's disease virus is used at 100 TCID50 per 0,025 ml; inactivated undiluted serum samples are mixed with an equal volume (0,025 ml) of virus suspension. The virus/serum mixtures are incubated for two hours at 37 °C in the microtitre plates before the appropriate cells are added. Cells are used at a concentration which forms a complete monolayer after

24 hours.

Controls: (i) virus infectivity assay, (ii) serum toxicity

controls, (iii) uninoculated cell culture controls,

(iv) reference antisera.

Interpretation:

The results of the neutralisation test and the titre of the virus used in the test are recorded after three to seven days incubation at 37 °C. Serum titres less than 1/2 (undiluted serum) are considered negative.

B. Any other test recognised in the framework of Decision 2008/185/EC (1).

Transmissible gastro-enteritis (TGE)

The serum neutralisation test shall be carried out according to the following protocol:

All sera are heat-inactivated at 56 °C for 30 minutes Serum:

before use.

Procedure: The constant virus-varying serum neutralisation test

on microtitre plates employs A72 (dog tumour) cells or other sensitive cell systems. TGE virus is used at 100 TCID50 per 0,025 ml; inactivated undiluted serum samples are mixed with an equal volume (0,025 ml) of virus suspension. The virus/serum mixtures are incubated for 30 to 60 minutes at 37 °C in the microtitre plates before the appropriate cells are added. Cells are used at a concentration which forms a complete monolayer after 24 hours.

Each cell receives 0,1 ml of cell suspension.

Controls: (i) virus infectivity assay, (ii) serum toxicity

controls, (iii) uninoculated cell culture controls,

(iv) reference antisera.

Interpretation: The results of the neutralisation test and the titre of

the virus used in the test are recorded after three to five days incubation at 37 °C. Serum titres less than 1/2 (final dilution) are considered negative. If undiluted serum samples are toxic to the tissue cultures, these sera may be diluted 1/2 before being used in the test. This is equivalent to 1/4 final dilution of serum. Serum titres of less than 1/4 (final dilution) are considered negative in these

cases.

Swine vesicular disease (SVD)

Tests for swine vesicular disease (SVD) shall be carried out according to Decision 2000/428/EC (2).

Classical swine fever (CSF)

Tests for classical swine fever (CSF) shall be carried out according to Decision 2002/106/EC (3).

The performance of tests for CSF must follow the guidelines set out in the relevant chapter of the OIE Manual of Diagnostic Tests and Vaccines for Terrestrial Animals.

⁽¹⁾ OJ L 59, 4.3.2008, p. 19.

⁽²⁾ OJ L 167, 7.7.2000, p. 22.

⁽³⁾ OJ L 39, 9.2.2002, p. 71.

▼C1

The evaluation of sensitivity and specificity of the serological test for CSF must be carried out in a national laboratory with a quality assurance scheme in place. Tests employed must be shown to recognise a range of weak and strong positive reference sera and allow detection of antibody in early phase and convalescence.

PART 7

Import and quarantine animal health conditions for animals imported into St. Pierre and Miquelon within a period of less than six months prior to introduction into the Union

(referred to in Article 6)

Animal species covered

-	Та	xon
ORDER	FAMILY	GENUS AND SPECIES
Artiodactyla	Camelidae	Camelus spp., Lama spp., Vicugna spp.

CHAPTER 1

Residence and quarantine

- Animals imported into St. Pierre and Miquelon must reside in an authorised quarantine station for a minimum period of 60 days before being dispatched for introduction into the Union. This period may be increased due to testing requirements for individual species. In addition the animals must comply with the following requirements:
 - (a) Separate consignments may enter the quarantine station. However, upon entry in the quarantine station all animals of the same species in the quarantine facility must be considered as a single group, and referred to as such. The quarantine period must commence for the whole group at the time when the last animal entered the quarantine facility.
 - (b) Within the quarantine station each specific group of animals must be maintained in isolation, with no direct or indirect contact with any other animals, including those from other consignments that may be present.

Each consignment must be kept in the approved quarantine station and protected from vector insects.

- (c) If, during the period of quarantine, the isolation of a group of animals is not maintained and contact is made with other animals, the quarantine period must begin again for the same duration as initially prescribed on entry into the quarantine station.
- (d) Animals to be introduced into the Union which pass through the quarantine station must be loaded and dispatched directly to the Union:
 - (i) without coming into contact with animals other than animals which fulfil the health conditions established for the introduction of the relevant category of animal into the Union;
 - (ii) segregated into consignments so that no consignment can came in contact with animals not eligible for importation into the Union;

- (iii) in transport vehicles or containers which have first been cleansed and disinfected with a disinfectant officially authorised in St. Pierre and Miquelon as effective in the control of the diseases referred to in Chapter 2 and which are so constructed that faeces, urine, litter or fodder cannot flow or fall out of the vehicle or container during transportation.
- 2. The quarantine premises must at least meet the minimum standards laid down in Annex B to Directive 91/496/EEC (¹), and the following conditions:
 - (a) they must be supervised by an official veterinarian;
 - (b) they must be situated at the centre of an area of at least 20 km in diameter in which, according to official findings, for at least 30 days prior to their use as a quarantine station there has been no case of foot-and-mouth disease;
 - (c) they must, before being used as a quarantine station, be cleansed and disinfected with a disinfectant officially authorised in St Pierre et Miquelon as effective in the control of the diseases referred to in Chapter 2;
 - (d) they must operate, taking into account their animal capacity:
 - (i) a facility dedicated exclusively for the quarantine of animals, including adequate housing to a suitable standard for the animals;
 - (ii) appropriate facilities, that:
 - are easy to thouroughly clean and disinfect,
 - include facilities for safe loading and unloading,
 - are able to fulfil all watering and feeding requirements for the animals,
 - allow any necessary veterinary treatment to be easily administered;
 - (iii) appropriate facilities for inspection and isolation;
 - (iv) appropriate equipment for cleaning and disinfecting rooms and transport vehicles;
 - (v) an appropriate storage area for fodder, litter and manure;
 - (vi) an appropriate system for collecting waste water;
 - (vii) an office for the official veterinarian;
 - (e) when operating, they must have sufficient veterinarians to carry out all duties:
 - (f) they must only admit animals that are individually identified so as to guarantee traceability. To this end, when animals are admitted the owner or the person in charge of the quarantine station must ensure that the animals are properly identified and accompanied by health certificates for the species and categories involved. In addition, the owner or the person in charge of the quarantine station must record on a register or in a data base, and retain for at least three years, the name of the owner, the origin of the animals in the consignment, the dates of entry and exit of the animals in the consignment, the identification number of the animals in the consignment and their place of destination;

(g) the competent authority must determine the procedure for official supervision of the quarantine station and must ensure that such supervision is carried out; this supervision must include regular inspections in order to ascertain that the requirements for approval continue to be fulfilled. In case of failure and suspension, the approval may only be restored when the competent authority is satisfied that the quarantine premises are in full compliance with all the conditions set out in points (a) to (g).

CHAPTER 2

Animal health tests

1. GENERAL REQUIREMENTS

The animals must be subjected to the following tests carried out on samples of blood taken, if not specified otherwise, not earlier than 21 days from the date of commencement of the isolation period.

The laboratory tests must be carried out in an approved laboratory in the Union and all laboratory tests and their results, vaccinations and treatments must be enclosed with the health certificate.

In order to keep animal interventions to a minimum, sampling, tests and any vaccinations must be grouped as far as is possible whilst respecting the minimum time intervals required by the testing protocols set out in Part 2 of this Chapter.

2. SPECIFIC REQUIREMENTS

2.1 CAMELIDAE

2.1.1 Tuberculosis

(a) Test to be used: comparative intradermal reaction test using Bovine purified protein derivative (PPD) and Avian PPD conforming to the standards for the manufacture of bovine and avian tuberculins as described in point 2.1.2 of Annex B of Directive 64/432/EEC.

The test must be executed in the area behind the shoulder (axillary region) following the technique described in point 2.2.4 of Annex B of Directive 64/432/EEC.

(b) Timing: the animals must be tested within two days from the date of arrival in the quarantine station and 42 days from the date of the first test.

(c) Interpretation of tests:

the reaction shall be considered:

- negative if the increased skin thickness is less than 2 mm.
- positive if the increased skin thickness is more than 4 mm.
- inconclusive if the increased skin thickness to the bovine PPD is between 2mm and 4 mm, or more than 4 mm but less then the reaction to the avian PPD.

(d) Options for action following testing:

If an animal presents a positive result to the intradermal-reaction to the bovine PPD, that animal shall be excluded from the group and the other animals shall be re-tested starting at least 42 days from the date of the first positive test was administered and this shall be considered as the first test described in (b).

If more than one animal of the group presents a positive result, the whole group shall be rejected for exportation to the Union.

If one or more animals of the same group present an inconclusive reaction, the whole group shall be re-tested starting at least 42 days from the date of the first test was administered and it shall be considered as the first test described in (b).

2.1.2 Brucellosis

(a) Test to be used:

- (i) Brucella abortus: Rose Bengal test (RBT) and Serum agglutination test (SAT) as described respectively in points 2.5 and 2.6 of Annex C to Directive 64/432/EEC. In the case of a positive result, a complement-fixation test shall be performed for confirmation as described in Part 6 of Annex I to Regulation (EU) No 206/2010.
- (ii) Brucella melitensis: RBT and SAT as described respectively in points 2.5 and 2.6 of Annex C to Directive 64/432/EEC. In the case of a positive result, a complement-fixation test following the method described in Annex C to Directive 91/68/EEC shall be performed for confirmation.
- (iii) Brucella ovis: Complement fixation test as described in Annex D to Directive 91/68/EEC
- (b) Timing: the animals have to be tested within two days from the date of their arrival in the quarantine station and 42 days from the date of the first test.

(c) Interpretation of tests:

A positive reaction to the tests shall be as defined in Annex C to Directive 64/432/EEC.

(d) Options for action following testing:

Animals tested positive to one of the tests shall be excluded from the group and the other animals shall be re-tested starting at least 42 days from the date the first positive test was performed: this shall be considered as the first test described in (b).

Only the animals that tested negative to two consecutive tests performed as described in (b) shall be allowed for the introduction to the Union.

2.1.3 Bluetongue and Epizootic haemorrhagic disease (EHD)

(a) Test to be used: agar gel immunodiffusion (AGID) test as described in Part 6 of Annex I to Regulation (EU) No 206/2010. In case of a positive reaction the animals shall be tested with competitive ELISA test as described in Part 6 of Annex I to Regulation (EU) No 206/2010 to discriminate between the two diseases.

(b) Timing:

The animals must be tested with negative result to two tests: the first within two days from the date of their arrival in the quarantine station and the second at least 21 days from date of the first test.

(c) Options for action following testing:

(i) Bluetongue

If one or more animals tested positive to the ELISA as described in Part 6 of Annex I to Regulation (EU) No 206/2010, the positive animal/animals shall be excluded from the group, and all the remaining animals in the group must be quarantined for 100 days starting from the date on which the samples for the positive test were collected. The group shall only be considered free of the bluetongue disease if regular checks carried out by official veterinarians throughout the duration of the quarantine period fail to reveal clinical symptoms of disease, and the quarantine station remains free of bluetongue vectors (*Culicoides*).

If a further animal presents clinical symptoms of bluetongue disease during the quarantine period as described in the first subparagraph, all the animals in the group shall be rejected for introduction into the Union.

(ii) Epizootic haemorrhagic disease (EHD)

If one or more animals tested positive reveal the presence of antibodies to the EHD virus during confirmatory ELISA testing, the animal(s) shall be considered positive and shall be excluded from the group, and the whole group shall be subject to repeat testing beginning at least 21 days from the date of the initial positive diagnosis and again at least 21 days from the date of the repeat test, both with negative results.

If any additional animals are tested positive during either or both of the two tests carried out for repeat testing, the whole group of animals shall be rejected for introduction into the Union.

2.1.4 Foot-and-Mouth Disease (FMD)

- (a) Test to be used: Diagnostic tests (probang and serology) using ELISA and (Virus Neutralisation) (VN) techniques in accordance with the Protocols described in Part 6 of Annex I to Regulation (EU) No 206/2010.
- (b) Timing: the animals shall be tested with negative results to two tests: the first within two days from the date of their arrival in the quarantine station and the second at least 42 days from the date of the first test.
- (c) Options for action following testing: If any animal tests positive for the FMD virus, then none of the animals present in the quarantine station shall be considered as eligible for introduction into the Union.

Note: Any detection of antibodies to structural or not structural proteins of FMD virus shall be considered as a result of previous infection of FMD irrespective of the vaccination status.

2.1.5 Rinderpest

- (a) Test to be used: The competitive ELISA test as described in the OIE Manual of Diagnostic Tests and Vaccines for Terrestrial Animals, latest version, is the prescribed test for international trade and is test of choice. Serum neutralisation test, or other recognised tests in accordance with the protocols described in relevant sections of the OIE manual may also be used.
- (b) Timing: the animals shall be tested twice: the first within two days from the date of their arrival in the quarantine station and the second at least 42 days from the date of the first test.
- (c) Options for action following testing: If any animal tests positive for the Rinderpest virus, then none of the animals present in the quarantine station shall be considered as eligible for introduction into the Union

2.1.6 Vesicular stomatitis

- (a) Test to be used: ELISA, virus neutralisation test, or other recognised test in accordance with the protocols described in the relevant sections of the OIE manual.
- (b) Timing: the animals shall be tested twice: the first within two days from the date of their arrival in the quarantine station and the second at least 42 days from the date of the first test.
- (c) Options for action following testing: If any animal tests positive for vesicular stomatitis virus, then none of the animals present in the quarantine station shall be considered as eligible for introduction into the Union.

2.1.7 Rift valley fever

- (a) Test to be used: ELISA, virus neutralisation test, or other recognised test in accordance with the protocols described in relevant sections of the OIE manual.
- (b) Timing: the animals shall be tested twice: the first within two days from the date of their arrival in the quarantine station and the second at least 42 days from the date of the first test.
- (c) Options for action following testing: If any animal displays evidence of exposure to rift valley fever agent, then none of the animals present in the quarantine station shall be considered as eligible for introduction into the Union.

2.1.8 Lumpy skin disease

(a) Test to be used: Serology using ELISA, virus neutralisation test, or other recognised test in accordance with the protocols described in relevant sections of the OIE manual.

- (b) Timing: the animals shall be tested twice: the first within two days from the date of their arrival in the quarantine station and the second at least 42 days from the date of the first test.
- (c) Options for action following testing: If any animal displays evidence of exposure to lumpy skin disease, then none of the animals present in the quarantine station shall be considered as eligible for introduction into the Union.

2.1.9 Crimean congo haemorrhagic fever

- (a) Test to be used: ELISA, virus neutralisation test, Immunofluorescence test or other recognised test.
- (b) Timing: the animals shall be tested twice: the first within two days from the date of their arrival in the quarantine station and the second at least 42 days from the date of the first test.
- (c) Options for action following testing: If any animal displays evidence of exposure to crimean congo haemorrhagic fever agent, then none of the animals present in the quarantine station shall be considered as eligible for introduction into the Union.

2.1.10 Surra (Trypanosoma evansi (T. evansi))

- (a) Test to be used: The parasitic agent can be identified in concentrated blood samples in accordance with the protocols described in relevant sections of the OIE manual.
- (b) Timing: the animals shall be tested twice: the first within two days from the date of their arrival in the quarantine station and the second at least 42 days from the date of the first test.
- (c) Options for action following testing: If T. evansi is detected in any animal in the consignment, then that animal shall be considered not eligible for introduction into the Union. The remaining animals of the group shall then undergo internal and external antiparasitic treatment using suitable agents that are effective against T. evansi.

2.1.11 Malignant catarrhal fever

- (a) Test to be used: Detection of viral DNA based on identification by immunofluorescence or immunocytochemistry using the protocols described in relevant sections of the OIE manual.
- (b) Timing: the animals shall be tested twice: the first within two days from the date of their arrival in the quarantine station and the second at least 42 days from the date of the first test.
- (c) Options for action following testing: If any animal displays evidence of exposure to MCF, then none of the animals present in the quarantine station shall be considered as eligible for introduction into the Union.

2.1.12 Rabies

Vaccination: Rabies vaccination may be carried out when requested by the Member State of destination and the animal shall be blood sampled and a serum neutralisation test for antibodies carried out.

- 2.1.13 *Enzootic bovine leucosis.* (only in the case where the animals are destined for an officially enzootic-bovine-leucosis free Member State or region, as referred to in Article 2(2)(k) of Directive 64/432/EEC)
 - (a) **Test to be used**: AGID or blocking ELISA, in accordance with the protocols described in the OIE manual, latest version.
 - (b) Timing: the animals shall be tested twice: the first within two days from the date of their arrival in the quarantine station and the second at least 42 days from the date of the first test.
 - (c) Options for action following testing: animals tested positive to the test described in (a) shall be excluded from the group of animals in the quarantine facility and the other animals shall be re-tested starting at least 21 days from the date of the first positive test was performed: this shall be considered as the first test described in (b).

Only the animals that tested negative to two consecutive tests performed as described in (b) shall be considered eligible for introduction into the Union.

FRESH MEAT

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PART 1
List of third countries, territories and parts thereof (1)

ISO code and name of third	Code of	Description of third country, territory or part thereof	Veterinary	certificate	Specific	Closing date (2)	Opening date (3)
country	Territory	Description of unite country, territory of part thereof	Model(s)	SG	conditions	Closing date (2)	Opening date (3)
1	2	3	4	5	6	7	8
AL – Albania	AL-0	Whole country	_				
AR – Argentina	AR-0	Whole country	EQU				
	AR-1	The Provinces of: Buenos Aires, Catamarca, Corrientes (except the departments of Berón de Astrada, Capital, Empedrado, General Paz, Itati, Mbucuruyá, San Cosme and San Luís del Palmar) Entre Ríos, La Rioja,	BOV	A	1		18 March 2005
		Mendoza, Misiones, Part of Neuquén (excluding territory included in AR-4), Part of Río Negro (excluding territory included in AR-4),	RUF	A	1		1 December 2007

1	2	3	4	5	6	7	8
		San Juan, San Luis, Santa Fe, Tucuman, Cordoba, La Pampa, Santiago del Estero,	RUW	A	1		1 August 2010
	AD 2	Chaco, Formosa, Jujuy and Salta, excluding the buffer area of 25 Km from the border with Bolivia and Paraguay that extends from the Santa Catalina District in the Province of Jujuy, to the Laishi District in the Province of Formosa					
	AR-2	Chubut, Santa Cruz and Tierra del Fuego	BOV, OVI, RUW, RUF				1 March 2002
	AR-3	Corrientes: the departments of Berón de Astrada, Capital, Empedrado, General Paz, Itati, Mbucuruyá, San Cosme and San Luís del Palmar	BOV RUF	A	1		1 December 2007
	AR-4	Part of Río Negro (except: in Avellaneda the zone located north of the Provincial road 7 and east of the Provincial road 250, in Conesa the zone located east of the Provincial road 2, in El Cuy the zone located north of the Provincial road 7 from its intersection with the Provincial road 66 to the border with the Department of Avellaneda, and in San Antonio the zone located east of the Provincial roads 250 and 2) Part of Neuquén (except in Confluencia the zone located east of the Provincial road 17, and in Picun Leufú the zone located east	BOV, OVI, RUW, RUF				1 August 2008
		of the Provincial road 17)					
AU – Australia	AU-0	Whole country	BOV, OVI, POR, EQU, RUF, RUW, SUF, SUW				

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1112								
	1	2	3	4	5	6	7	8
	BA – Bosnia and Herzegovina	BA-0	Whole country	_				
	BH – Bahrain	BH-0	Whole country					
	BR – Brazil	BR-0	Whole country	EQU				
		BR-1	State of Minas Gerais					
			State of Espírito Santo;					
			State of Goiás;					
			State of Mato Grosso					
			State of Rio Grande Do Sul, State of Mato Grosso Do Sul (except for the designated high surveillance zone of 15 Km from the external borders in the municipalities of Porto Murtinho, Caracol, Bela Vista, Antônio João, Ponta Porã, Aral Moreira, Coronel Sapucaia, Paranhos, Sete Quedas, Japorã, and Mundo Novo and the designated high surveillance zone in the municipalities of Corumbá and Ladário).	BOV	A and H	1		1 December 2008
		BR-2	State of Santa Catarina	BOV	A and H	1		31 January 2008
		BR-3	States of Paraná and São Paulo	BOV	A and H	1		1 August 2008
<u>M4</u>								
	BW – Botswana	BW-0	Whole country	EQU, EQW				
		BW-1	The veterinary disease control zones 3c, 4b, 5, 6, 8, 9 and 18	BOV, OVI, RUF, RUW	F	1	11 May 2011	1 December 2007
		BW-2	The veterinary disease control zones 10, 11, 13 and 14	BOV, OVI, RUF, RUW	F	1		7 March 2002
		BW-3	The veterinary disease control zone 12	BOV, OVI, RUF, RUW	F	1	20 October 2008	20 January 2009
		BW-4	The veterinary disease control zone 4a, except the intensive surveillance buffer zone of 10 km along the boundary with the foot-and-mouth disease vaccination zone and wildlife management areas	BOV	F	1		18 February 2011

1	2	3	4	5	6	7	8
BY – Belarus	BY-0	Whole country	_				
BZ – Belize	BZ-0	Whole country	BOV, EQU				
CA – Canada	CA-0	Whole country	BOV, OVI, POR, EQU, SUF, SUW, RUF, RUW	G			
CH – Switzerland	CH-0	Whole country	*				
CL – Chile	CL-0	Whole country	BOV, OVI, POR, EQU, RUF, RUW, SUF				
CN – China	CN-0	Whole country	_				
CO – Colombia	CO-0	Whole country	EQU				
CR – Costa Rica	CR-0	Whole country	BOV, EQU				
CU – Cuba	CU-0	Whole country	BOV, EQU				
DZ – Algeria	DZ-0	Whole country	_				
ET – Ethiopia	ET-0	Whole country	_				
FK – Falkland Islands	FK-0	Whole country	BOV, OVI, EQU				
GL – Greenland	GL-0	Whole country	BOV, OVI, EQU, RUF, RUW				
GT – Guatemala	GT-0	Whole country	BOV, EQU				

1	2	3	4	5	6	7	8
HK – Hong Kong	HK-0	Whole country	_				
HN – Honduras	HN-0	Whole country	BOV, EQU				
HR – Croatia	HR-0	Whole country	BOV, OVI, EQU, RUF, RUW				
IL – Israel	IL-0	Whole country	_				
IN – India	IN-0	Whole country	_				
IS – Iceland	IS-0	Whole country	BOV, OVI, EQU, RUF, RUW				
KE – Kenya	KE-0	Whole country	_				
MA – Morocco	MA-0	Whole country	EQU				
ME – Montenegro	ME-0	Whole country	BOV, OVI, EQU				
MG – Madagascar	MG-0	Whole country	_				
MK – Former Yugoslav Republic of Macedonia (4)	MK-0	Whole country	OVI, EQU				
MU – Mauritius	MU-0	Whole country	_				
MX – Mexico	MX-0	Whole country	BOV, EQU				
NA – Namibia	NA-0	Whole country	EQU, EQW				
	NA-1	South of the cordon fences which extend from Palgrave Point in the west to Gam in the east	BOV, OVI,RUF, RUW	F and J	1		
NC – New Caledonia	NC-0	Whole country	BOV, RUF, RUW				

1	2	3	4	5	6	7	8
NI – Nicaragua	NI-0	Whole country	_				
NZ – New Zealand	NZ-0	Whole country	BOV, OVI, POR, EQU, RUF, RUW, SUF, SUW				
PA – Panama	PA-0	Whole country	BOV, EQU				
PY – Paraguay	PY-0	Whole country	EQU				
	PY-1	Whole country except for the designated high surveillance zone of 15 Km from the external borders	BOV	A	1		1 August 2008
RS – Serbia (⁵)	RS-0	Whole country	BOV, OVI, EQU				
RU – Russia	RU-0	Whole country	_				
	RU-1	Region of Murmansk, Yamolo-Nenets autonomous area	RUF				
SV – El Salvador	SV-0	Whole country	_				
SZ – Swaziland	SZ-0	Whole country	EQU, EQW				
	SZ-1	Area west of the 'red line' fences which extends northwards from the river Usutu to the frontier with South Africa west of Nkalashane,	BOV, RUF, RUW	F	1		
	SZ-2	The veterinary foot and mouth disease surveillance and vaccination control areas as gazetted as a Statutory Instrument under legal notice number 51 of 2001	BOV, RUF, RUW	F	1		4 August 2003

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	1	2	3	4	5	6	7	8
TH – T	Thailand	TH-0	Whole country					
TN – Tunisia TN-0		TN-0	Whole country	_				
TR – T	`urkey	TR-0	Whole country	_				
		TR-1	The provinces of Amasya, Ankara, Aydin, Balikesir, Bursa, Cankiri, Corum, Denizli, Izmir, Kastamonu, Kutahya, Manisa, Usak, Yozgat and Kirikkale	EQU				
UA – U	Jkraine	UA-0	Whole country	_				
US – U	United States	US-0	Whole country	BOV, OVI, POR, EQU,SUF, SUW, RUF, RUW	G			
UY – U	Jruguay	UY-0	Whole country	EQU				
				BOV,	A	1		1 November 2001
				OVI	A	1		
<u></u>								
ZA – S	South Africa	ZA-0	Whole country	EQU, EQW				
		ZA-1	The whole country except: — the part of the foot-and-mouth disease control area situated in the veterinary regions of Mpumalanga and Northern provinces, in the district of Ingwavuma of the veterinary region of Natal and in the border area with Botswana east of longitude 28°, and — the district of Camperdown, in the province of KwaZulu-Natal.	BOV, OVI, RUF, RUW	F	1	11 February 2011	

1	2	3	4	5	6	7	8
ZW – Zimbabwe	ZW-0	Whole country	_				

Footnotes:

- (1) Without prejudice to specific certification requirements provided for in Union agreements with third countries.
- (2) Meat from animals slaughtered on or before the date set out in column 7 may be imported into the Union for 90 days from that date. Consignments carried on vessels on the high seas may be imported into the Union if certified before the date set out in column 7 for 40 days from that date. (N.B.: no date in column 7 means that there are no time restrictions).
- (3) Only meat from animals slaughtered on or after the date set out in column 8 may be imported into the Union (no date in column 8 means that there are no time restrictions).
- (4) The former Yugoslav Republic of Macedonia; provisional code that does not prejudge in any way the definitive nomenclature for this country, which will be agreed following the conclusion of negotiations currently taking place on this subject in the United Nations.
- (5) Not including Kosovo which is at present under international administration pursuant to United Nations Security Council Resolution 1244 of 10 June 1999
- * = Requirements as in accordance with the Agreement between the European Community and the Swiss Confederation on trade in agricultural products (OJ L 114, 30.4.2002, p. 132).
- = No certificates are laid down and fresh meat imports are prohibited (except for those species where indicated in the line comprising the entry for the whole country).

'1' Category restrictions:

No offal is authorised for introduction into the Union (except, in the case of bovine species, diaphragm and masseter muscles).

PART 2

Models of veterinary certificates

Model(s):

'BOV': Model of veterinary certificate for fresh meat, including minced meat, of domestic bovine animals (including *Bison* and *Bubalus*

species and their cross-breeds).

'OVI': Model of veterinary certificate for fresh meat, including minced

meat, of domestic ovine animals (Ovis aries) and domestic caprine

animals (Capra hircus).

'POR': Model of veterinary certificate for fresh meat, including minced

meat, of domestic porcine animals (Sus scrofa).

'EQU': Model of veterinary certificate for fresh meat, excluding minced

meat, of domestic solipeds (Equus caballus, Equus asinus and

their cross-breeds).

'RUF': Model of veterinary certificate for fresh meat, excluding offal and

minced meat, of farmed non-domestic animals of the order Artio-dactyla (excluding bovine animals (including *Bison* and *Bubalus* species and their cross-breeds), *Ovis aries*, *Capra hircus*, Suidae and Tayassuidae), and of the families Rhinocerotidae and

Elephantidae.

'RUW': Model of veterinary certificate for fresh meat, excluding offal and

minced meat, of wild non-domestic animals of the order Artiodactyla (excluding bovine animals (including *Bison* and *Bubalus* species and their cross-breeds), *Ovis aries, Capra hircus*, Suidae and Tayassuidae), and of the families Rhinocerotidae and Elephantidae.

'SUF': Model of veterinary certificate for fresh meat, excluding offal and

minced meat, of farmed non-domestic animals belonging to the

Suidae, Tayassuidae, or Tapiridae families.

'SUW': Model of veterinary certificate for fresh meat, excluding offal and

minced meat, of wild non-domestic animals belonging to the Suidae,

Tayassuidae, or Tapiridae families.

'EQW': Model of veterinary certificate for fresh meat, excluding offal and

minced meat, of wild solipeds belonging to the subgenus Hippotigris

(zebra).

SG (Supplementary guarantees)

'A': guarantees regarding the maturation, pH measurement and boning of

fresh meat, excluding offal, certified according to the models of veterinary certificates BOV (point II.2.6), OVI (point II.2.6), RUF

(point II.2.7) and RUW (point II.2.4).

'C': guarantees regarding the laboratory test for classical-swine-fever in the carcases from which fresh meat was obtained, certified according

to the model of veterinary certificate SUW (point II.2.3 B).

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- 'D': guarantees regarding swill feed on holding(s) of animals from which fresh meat certified was obtained according to the model of veterinary certificate POR (point II.2.3 d).
- 'E': guarantees regarding tuberculosis test in the animals from where fresh meat certified was obtained, according to the model of veterinary certificate BOV (point II.2.4 d).
- 'F': guarantees regarding the maturation and de-boning of fresh meat, excluding offal, certified according to the models of veterinary certificates BOV (point II.2.6), OVI (point II.2.6), RUF (point II.2.6) and RUW (point II.2.7).
- 'G': guarantees regarding 1, exclusion of offals and spinal cord; and 2, testing and origin of cervid animals in relation to chronic wasting disease as referred to in the models of veterinary certificates RUF (point II.1.7) and RUW (point II.1.8).
- 'H': supplementary guarantees required for Brazil. Concerning vaccination programmes, as the State of Santa Catarina in Brazil does not vaccinate against foot and mouth disease, the reference to a vaccination programme is not applicable for meat coming from animals originating and slaughtered in that State.
- 'J': guarantees regarding the movement of bovine, ovine and caprine animals from holdings to the slaughterhouse, which allow them to pass via an assembly centre (including markets) before being transported directly to slaughter.

Model BOV

ου	NTRY							Veterinary of	ertificate to El	
	l.1.	Consignor		I.2. Certi	ficate refe	erence No	1.2	.a.		
		Name		I.3. Cent	ral compe	etent authority				
		Address								
Ę		Tel.		I.4. Loca	l compete	ent authority				
Part I: Details of dispatched consignment	1.5.	Consignee		1.6.						
onsig		Name								
გ გ		Address								
atch		Postal code								
disb		Tel.								
s of	1.7.	Country of origin ISO code I.8. Region	n of origin Code		ntry of ination	ISO code	I.10. Reg	jion of tination	Code	
Detai				4001			400			
=======================================	l.11.	Place of origin	-	I.12.						
Pa		Name Approval nu	mber			_				
		Address								
	112	Place of loading		I.14. Date of departure						
	1.15.	r lace of loading		livii Bato	or dopare	u. 0				
	l.15.	Means of transport		I.16. Entry	BIP in E	U				
			Railway wagon 🔲							
		Road vehicle Other Identification		l.17.						
		Documentary references								
	I.18.	Description of commodity		I.19. Commodity code (HS code)						
							I.20. Quar	ntity		
	1.21.	Temperature of product					I.22. Numb	ber of packa	ges	
		Ambient	7	Frozen [٦					
	1.23.	Seal/Container No			<u> </u>		I.24. T ype	of packagin	g	
	1.25.	Commodities certified for:								
		Human consumption								
	1.26.			127 For it	mnort or r	admission int	^ EU		1	
	1.20.			1.21. FUI II	mport or a	admission int	0 20	_	ı	
	1.28.	Identification of the commodities		1						
		Species Nature of commodity	Treatment A	Approval number of establishments Number of Net packages weight						
					- •					

COUNTRY Model BOV Health information II.b. II.a. Certificate reference number 11.1. **Public Health Attestation** I, the undersigned official veterinarian declare that I am aware of the relevant requirements of Regulations (EC) No 178/2002, (EC) No 852/2004, (EC) No 853/2004, (EC) No 853/2004, (EC) No 854/2004 and (EC) No 999/2001 and certify that the meat of domestic bovine animals described in Part I was produced in accordance with those requirements, in particular that: Certification II.1.1. the [meat] [minced meat] (1) comes from (an) establishment(s) implementing a programme based on the HACCP principles in accordance with Regulation (EC) No 852/2004; II.1.2. the meat has been obtained in compliance with Section I of Annex III to Regulation (EC) No 853/2004; Part (1) II.1.3. [the minced meat has been produced in compliance with Section V of Annex III to Regulation (EC) No 853/2004 and frozen to an internal temperature of not more than – 18 °C;] II.1.4. the meat has been found fit for human consumption following ante and post-mortem inspections carried out in accordance with Chapter II of Section I and Chapters I and IX of Section IV of Annex I to Regulation (EC) No 854/2004; II.1.5. (1) either [the carcass or parts of the carcass have been marked with a health mark in accordance with Chapter III of Section I of Annex I to Regulation (EC) No 854/2004;] [the packages of [meat] [minced meat] (1) have been marked with an identification mark in accordance with Section I of Annex II to Regulation (EC) No 853/2004;] (1) or II.1.6. the [meat] [minced meat] (1) satisfies the relevant criteria set out in Regulation (EC) No 2073/2005 on microbiological criteria for II.1.7. the guarantees covering live animals and products thereof provided by the residue plans submitted in accordance with Directive 96/23/EC, and in particular Article 29 thereof, are fulfilled; II.1.8. the [meat] [minced meat] (¹) has been stored and transported in accordance with the relevant requirements of Sections I and V respectively of Annex III to Regulation (EC) No 853/2004; II.1.9. with regard to bovine spongiform encephalopathy (BSE): (1) either [II.1.9.1. for imports from a country or a region with a negligible BSE risk and listed as such in Decision 2007/453/EC: (a) the country or region is classified in accordance with Article 5(2) of Regulation (EC) No 999/2001 as a country or region posing a negligible BSE risk; (b) the animals from which the bovine meat or minced meat was derived were born, continuously reared and slaughtered in a country with a negligible BSE risk (¹³); $(^1)$ [(c) if in the country or region there have been BSE indigenous cases: (1) either [the animals were born after the date from which the ban on the feeding of ruminants with meatand-bone meal and greaves derived from ruminants had been enforced.] [the bovine meat or minced meat does not contain and is not derived from specified risk material (1) or as defined in Annex V to Regulation (EC) No 999/2001, or mechanically separated meat obtained from bones of bovine animals.]]] [II.1.9.2. for imports from a country or a region with a controlled BSE risk and listed as such in Decision 2007/453/EC: (1) or

> (a) the country or region is classified in accordance with Article 5(2) of Regulation (EC) No 999/2001 as a country or region posing a controlled BSE risk;

COUNT	'RY	Model BOV
II.	Health information	II.a. Certificate reference number II.b.
		(b) the animals from which the bovine meat or minced meat was 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) either [(c) the bovine meat or minced meat does not contain and is not derived from specified risk material as defined in Annex V to Regulation (EC) No 999/2001, or mechanically separated meat obtained from bones of bovine animals.]
		(¹) or [(c) the carcasses, half carcasses or half carcasses cut into no more than three wholesale cuts, and quarters contain no specified risk material other than the vertebral column, including dorsal root ganglia. The carcasses or wholesale cuts of carcasses of bovine animals containing vertebral column have been identified by a blue stripe on the label referred to in Regulation (EC) No 1760/2000. (³)]]
	(¹) or [II.1.9.3.	for imports from a country or a region which has not been categorised in accordance with Article 5(2) of Regulation (EC) No 999/2001 or has been categorised as a country or region with undetermined BSE risk and listed as such in Decision 2007/453/EC:
		(a) the country or region has not been categorised in accordance with Article 5(2) of Regulation (EC) No 999/2001 or has been categorised as a country or region with undetermined BSE risk;
		(b) the animals from which the bovine meat or minced meat was derived have not been fed meat-and-bone meal or greaves derived from ruminants;
		(c) the animals from which the bovine meat or minced meat was 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) eithe	er [(d) the bovine meat or minced meat was not derived from:
		(i) specified risk material as defined in Annex V to Regulation (EC) No 999/2001;
		(ii) nervous and lymphatic tissues exposed during the deboning process;
		(iii) mechanically separated meat obtained from bones of bovine animals.]
	(¹) or	[(d) the carcasses, half carcasses or half carcasses cut into no more than three wholesale cuts, and quarters contain no specified risk material other than the vertebral column, including dorsal root ganglia. The carcasses or wholesale cuts of carcasses of bovine animals containing vertebral column have been identified by a blue stripe on the label referred to in Regulation (EC) No 1760/2000. (3)]]
	Parl	iffils the requirements of Regulation (EC) No 1688/2005 implementing Regulation (EC) No 853/2004 of the European lament and of the Council as regards special guarantees concerning Salmonella for consignments to Finland and eden of certain meat and eggs;]
II.2.	Animal Health att	estation
	I, the undersigned	l official veterinarian, hereby certify, that the fresh meat described in Part I:
	II.2.1. has be	en obtained in the territory/ies with code:(2) which, at the date of issuing this certificate:
		as been free for 12 months from rinderpest, and during the same period no vaccination against this disease has taken ace, and
		as been free for 12 months from foot-and-mouth disease, and during the same period no vaccination against this disease as taken place;]
		is been considered free from foot-and-mouth disease since (dd/mm/yyyy), without having had cases/outbreaks terwards, and authorised to export this meat by Commission Regulation (EU) No/, of (dd/mm/yyyy);]

▼ M1

COUNTRY Model BOV Health information II.a. Certificate reference number II.b. (1) (5) or [(b) vaccination programmes against foot-and-mouth disease are being officially carried out and controlled in domestic bovine (1) (6) or [(b) has a systematic vaccination programme against foot and mouth disease and from herds where the efficacy of this vaccination programme is controlled by the competent veterinary authority through a regular serological surveillance indicating adequate antibody levels and which also demonstrates the absence of foot and mouth virus circulation;] (1) (6) or [(b) has been free for 12 months from foot-and-mouth disease, and during the same period no vaccination against this disease has taken place and is controlled by the competent veterinary authority through a regular surveillance demonstrating the absence of foot and mouth infection;] 11.2.2. has been obtained from animals that: (1) either [have remained in the territory described under point II.2.1 since birth, or for at least the last three months before slaughter;] (1) or (1) or Member State:1. 11.2.3. has been obtained from animals coming from holdings in which: (a) None of the animals present therein have been vaccinated against [foot-and-mouth disease or] (7) rinderpest, and [(b) in these holdings, and in the holdings situated in their vicinity within 10 km, there has been no case/outbreak of foot-and-(1) either mouth disease or rinderpest during the previous 30 days,] (1) (8) or (b there is no official restriction for animal health reasons and where, in these holdings and in the holdings situated in their vicinity within 25 km, there has been no case/outbreak of foot-and-mouth disease or rinderpest during the previous 60 days, and, (c) they have remained for at least 40 days before direct dispatch to the slaughterhouse;] (1) (14) or (c) they have remained for at least 40 days before passing through one assembly centre approved by the competent veterinary authority without coming into contact with animals of a different health status prior to subsequently going directly to a slaughterhouse;] (1) (9) or [(b) there is no official restriction for animal health reasons and where, in these holdings and in the holdings situated in their vicinity within 10 km, there has been no case/outbreak of foot-and-mouth disease or rinderpest during the previous 12 (c) they have remained for at least 40 days before direct dispatch to the slaughterhouse;] $(^1)(^6)$ (d) animals have not been introduced during the last 3 months from areas not approved by the EU: (e) animals are identified and registered in the national System of Identification and Certification of Origin for bovine animals: (f) the holdings in question are listed as approved holdings, following a favourable competent authorities' inspection and official report, in TRACES (10) and inspections are regularly carried out by the competent authorities to ensure that the relevant requirements provided for in Regulation (EU) No 206/2010 are respected.] II.2.4. has been obtained from animals which:

(a) have been transported from their holdings in vehicles, cleaned and disinfected before loading, to an approved slaughterhouse without contact with other animals which did not comply with the conditions referred to in point II.2.1, II.2.2 and II.2.3,

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COUNTRY Model BOV Health information II.a. Certificate reference number II.b. (b) at the slaughterhouse, have passed ante-mortem health inspection during the 24 hours before slaughter and, in particular, have shown no evidence of the diseases referred to in point II.2.1, (1) (12) [(d) have reacted negatively to an official intra-dermal tuberculosis test carried out within 3 months before slaughter;] (1) (6) [(e) at the slaughterhouse have been kept prior to slaughter completely separate from animals the meat of which is not intended for the Union1 II.2.5. has been obtained in an establishment around which, within a radius of 10 km, there has been no case/outbreak of the diseases referred to in point II.2.1 during the previous 30 days or, in the event of a case/outbreak of disease, the preparation of meat for importation to the Union has been authorised only after slaughter of all animals present, removal of all meat, and the total cleaning and disinfection of the establishment under the control of an official veterinarian; 11.2.6. [has been obtained and prepared without contact with other meats not complying with the conditions required in this certificate.] (1) either [contains [boneless meat] [and] [minced meat] $(^1)$, obtained only from de-boned meat other than offal that was obtained from carcasses in which the main accessible lymphatic glands have been removed, which have been submitted to maturation at a temperature above + 2 °C for at least 24 hours before the bones were removed and in which the pH value of the meat was below 6.0 when tested electronically in the middle of the longissimus-dorsi muscle after (1) (8) or maturation and before de-boning, and has been kept strictly separate from meat not conforming to the requirements referred to in this certificate during all stages of its production, de-boning and storage until it has been packed in boxes or cartons for further storage in dedicated areas.1 [contains [boneless meat] [and] [minced meat] (1), obtained only from de-boned meat other than offal that was obtained (1) (9) or from carcasses in which the main accessible lymphatic glands have been removed, which have been submitted to maturation at a temperature above + 2 °C for at least 24 hours before the bones were removed, and has been kept strictly separate from meat not conforming to the requirements referred to in this certificate during all stages of its production, de-boning and storage until it has been packed in boxes or cartons for further storage in

II.3. Animal welfare attestation

dedicated areas.1

I, the undersigned official veterinarian, hereby certify, that the fresh meat described in Part I derives from animals which have been treated in the slaughterhouse before and at the time of slaughter or killing in accordance with the relevant provisions of Union legislation.

Notes

This certificate is meant for fresh meat, including minced meat, of domestic bovine animals (including *Bison* and *Bubalus* species and their cross-breeds).

Fresh meat means all animal parts fit for human consumption whether fresh, chilled or frozen.

Part I

- Box reference I.8: Provide the code of territory as appearing in Part 1 of Annex II to Regulation (EU) No 206/2010.
- Box reference I.11: Place of origin: name and address of the dispatch establishment.
- Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor must inform the BIP of entry into the Union.
- Box reference I.19: Use the appropriate HS code: 02.01, 02.02, 02.06 or 05.04. In addition, for those territories of origin without the entry "A" or "F" in column 5 "SG" of Part 1 of Annex II to Regulation (EU) No 206/2010, the HS code 15.02 may also be used when appropriate.

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cou	UNTRY		Model BOV							
II.	Health information	II.a. Certificate reference number	II.b.							
_	Box reference I.20: Indicate total gross weight and total net weight.									
-	Box reference I.23: For containers or boxes, the container number a	and the seal number (if applicable) m	ust be included.							
-	Box reference I.28: Nature of commodity: Indicate "carcass-whole",	"carcass-side", "carcass-quarters", "cu	ts", "offal" or "minced meat".							
	Minced meat is deboned meat that has been minced into fragments and that must have been prepared exclusively from striated muscle (including the adjoining fatty tissues) except heart muscle.									
-	Box reference I.28: Treatment type: If appropriate, indicate "debone	d"; "bone in"; "matured"								
Par	t II:									
(¹)	Keep as appropriate.									
(2)	Code of the territory as it appears in Part 1 of Annex II to Regulation	on (EU) No 206/2010.								
(3)	The number of bovine carcasses or wholesale cuts of carcasses, from which removal of the vertebral column is required as well as the number where removal of the vertebral column is not required must be added to the common veterinary entry document referred to in Article 2 (1) of Regulation (EC) No 136/2004.									
(4)	Delete if the consignment is not intended for introduction into Finlar	nd or Sweden.								
(5)	Only matured de-boned meat fulfilling the supplementary guarantees	s referred to in footnote (8).								
(⁶)	Supplementary guarantees regarding import of matured de-boned meat to be provided when required in column 5 "SG" of Part 1 of Annex II to Regulation (EU) No 206/2010 with the entry "H".									
(7)	Delete when the exporting country carries out vaccination against foot-and-mouth disease with serotypes A, O or C, and this country is allowed to import into the Union matured de-boned meat which fulfils the supplementary guarantees described, in footnote (8).									
(8)	Supplementary guarantees regarding meats from matured de-boned meat to be provided when required in column 5 "SG" of Part 1 of Annex II to Regulation (EU) No 206/2010, with the entry "A".									
(9)	Supplementary guarantees regarding meats from matured de-boned II to Regulation (EU) No 206/2010, with the entry "F". The matured de days after the date of slaughter of the animals.									
(10)	The list of approved holdings provided by the competent authority authority. The Commission will ensure that this list of approved holding integrated computerised veterinary system (TRACES).									
(11)	Date or dates of slaughter. Imports of this meat shall not be allow authorisation for importation into the Union of the third country, terri where restrictive measures have been adopted by the Union again.	itory or part thereof referred to in box	es I.7 and I.8, or during a period							
(12)	Supplementary guarantees concerning tuberculosis test, to be provid (EU) No 206/2010, with the entry "E". Intra-dermal tuberculosis test to 64/432/EEC.									
(13)	List of countries in the Annex to Decision 2007/453/EC.									
(14)	Alternative guarantee may be provided when allowed for by the entry "J" in column 5 "SG" of Part 1 of Annex II to Regulation (EU) No 206/2010.									
Offi	cial veterinarian									
	Name (in capital letters):	Qualifica	tion and title:							
	Date:	Signature	o:'							
	Stamp:									

▼<u>M1</u>

Model OVI

100	NTRY										Veterinary cert	ificate to EU	
	l.1.	Consignor				1.2.	Certificat	e refe	erence No		1.2.a.		
		Address					I.3. Central competent authority						
ŧ							Local co	mpete	ent authority	'			
au d	l.5.	Consignee				1.6.							
isic		Name											
핥		Address											
gc		Postal code					_						
disp		Tel.	Tel.										
Part I: Details of dispatched consignment	1.7.	Country of origin	ISO code	I.8. Region of origin	Code	1.9.	Country destination		ISO code	I.10. R	Region of destinatio	n Code	
<u>ٿ</u> ت	1.11.	.11. Place of origin					1.12.						
Pa		Name Approval number Address											
	I.13. Place of loading					I.14. Date of departure							
	I.15. Means of transport					I.16. Entry BIP in EU							
		Aeroplane	Ship 🗌										
		Road vehicle Other Identification				1.17.							
		Documentary references											
	l.18.	Description of commodity						l.19.	Commodity	/ code (H	HS code)		
										1.20. Qu	uantity		
	1.21.	Temperature of produ	Femperature of product					I.22. Number of packages					
		Ambient 🗌		Chilled		Frozen							
	1.23.	Seal/Container No				I.24. Type of packaging							
	1.25.	Commodities certified	I for:							•			
		Human consumption											
	1.26.					1.27.	For impo	rt or	admission i	nto EU			
	1.28.	3. Identification of the commodities											
		Species (scientific name)	Nature commod		Abatto		al numbe Cutting		stablishmer It Co	ts d store	Number of packages	Net weight	

▼ M1

COUNTRY Model OVI

II. Health information II.a. Certificate reference number II.b.

1.1. Public Health Attestation

I, the undersigned official veterinarian, declare that I am aware of the relevant requirements of Regulations (EC) No 178/2002, (EC) No 852/2004, (EC) No 853/2004, (EC) No 854/2004 and (EC) No 999/2001 and certify that the meat of domestic ovine and caprine animals described in Part I was produced in accordance with those requirements, in particular that:

- II.1.1. the [meat] [minced meat] (1) comes from (an) establishment(s) implementing a programme based on the HACCP principles in accordance with Regulation (EC) No 852/2004;
- (1) II.1.2. the meat has been obtained in compliance with the conditions set out in Section I of Annex III to Regulation (EC) No 853/2004;
- (1) II.1.3. [the minced meat has been produced in compliance with Section V of Annex III to Regulation (EC) No 853/2004 and frozen to an internal temperature of not more than 18 °C;]
 - II.1.4. the meat has been found fit for human consumption following ante and post-mortem inspections carried out in accordance with Chapter II of Section I and Chapters II and IX of Section IV of Annex I to Regulation (EC) No 854/2004;
 - II.1.5. (1) either [the carcass or parts of the carcass have been marked with a health mark in accordance with Chapter III of Section I of Annex I to Regulation (EC) No 854/2004;]
 - (1) or [the packages of [meat] [minced meat] (1) have been marked with an identification mark in accordance with Section I of Annex II to Regulation (EC) No 853/2004;]
 - II.1.6. the [meat] [minced meat] (1) satisfies the relevant criteria set out in Regulation (EC) No 2073/2005 on microbiological criteria for foodstuffs:
 - II.1.7. the guarantees covering live animals and products thereof provided by the residue plans submitted in accordance with Directive 96/23/EC, and in particular Article 29 thereof, are fulfilled;
 - II.1.8. the [meat] [minced meat] (1) has been stored and transported in accordance with the relevant requirements of Sections I and V respectively of Annex III to Regulation (EC) No 853/2004;
 - II.1.9. with regard to bovine spongiform encephalopathy (BSE):
- (1) either [II.1.9.1. for imports from a country or a region with a negligible BSE risk and listed as such in Decision 2007/453/EC:
 - (a) the country or region is classified in accordance with Article 5(2) of Regulation (EC) No 999/2001 as a country or region posing a negligible BSE risk;
 - (b) the animals from which the meat or minced meat was derived were born, continuously reared and slaughtered in a country with negligible BSE risk; (2)
 - (1) [(c) if in the country or region there have been BSE indigenous cases:
 - (1) either [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 had been enforced.]
 - (1) or [the meat or minced meat does not contain and is not derived from specified risk material as defined in Annex V to Regulation (EC) No 999/2001, or mechanically separated meat obtained from bones of domestic ovine or caprine animals.]]]
- (1) or [II.1.9.2. for imports from a country or a region with a controlled BSE risk and listed as such in Decision 2007/453/EC:
 - (a) the country or region is classified in accordance with Article 5(2) of Regulation (EC) No 999/2001 as a country or region posing a controlled BSE risk;
 - (b) animals from which the meat or minced meat was 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;

Part II: Certification

COUN	TRY				Model OVI		
II.	Health i	information		II.a. Certificate reference number	II.b		
		(¹) either	[(c) the meat or minced meat does not conta Regulation (EC) No 999/2001, or mecha animals.]				
		(¹) or	[(c) the carcasses, half carcasses or half ca no specified risk material other than the				
	(¹) or	[II.1.9.3.	for imports from a country or a region whic (EC) No 999/2001 or has been categorised Decision 2007/453/EC:				
		 (a) the country or region has not been categorised in accordance with Article 5(2) of Regulation (EC) No 999/2001 has been categorised as a country or region with undetermined BSE risk; 					
	(b) the animals from which the meat or minced meat was derived have not been fed meat-and-bone meal or gre derived from ruminants;						
	(c) the animals from which the meat or minced meat was derived have not been slaughtered after stunning by m of gas injected into the cranial cavity or killed by the same method or slaughtered by laceration after stunni central nervous tissue by means of an elongated rod-shaped instrument introduced into the cranial cavity;						
		(1) either	[(d) the meat or minced meat was not deriv	ed from:			
			(i) specified risk material as defined in	Annex V to Regulation (EC) No 999/2	2001;		
			(ii) nervous and lymphatic tissues expo	sed during the deboning process;			
	(iii) mechanically separated meat obtained from bones of domestic ovine or caprine animal						
		(¹) or	[(d) the carcasses, half carcasses or half ca no specified risk material other than the				
II.2.	Animal	Health atte	estation				
	I, the u	ndersigned	official veterinarian, hereby certify, that the fi	resh meat described in Part I:			
	II.2.1.	has been	obtained in the territory/ies with code:	(3) which, at the date of iss	uing this certificate:		
		(a) has be	een free for 12 months from rinderpest, and d	during the same period no vaccination against this disease has taken place,			
	(¹) eithei		neen free for 12 months from foot-and-mouth aken place;]	disease, and during the same period	no vaccination against this disease		
	(¹) or	break	neen considered free from foot-and-mouth dis is afterwards, and authorised to export this r nm/yyyy);]				
	(¹) (⁴) or	(b) vaccii anima	nation programmes against foot-and-mouth oals;]	disease are being officially carried out	and controlled in domestic bovine		
	II.2.2.	has been	obtained from animals that:				
		(1) either	[have remained in the territory described slaughter;]	under point II.2.1 since birth, or for at	least the last three months before		
		(¹) or	[have been introduced onterritory with code (3) that at that date				
		(¹) or	[have been introduced on	(dd/mm/yyyy) into the territory describ	ped under point II.2.1, from the EU		

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cou	OUNTRY Model OVI								
II.	Healt	n information	II.a. Certificate reference number	II.b.					
	II.2.3.	has been obtained from animals coming from holdings:							
		(a) in which none of the animals present therein have be	een vaccinated against [foot-and-mouth	n disease or] (5) rinderpest,					
	(b) not subject to prohibition as a result of an outbreak of ovine or caprine brucellosis during the previous six weeks, and								
	(1) either [(c) in and around which, in an area of 10 km radius, there has been no case/outbreak of foot-and-mouth disease or rinderpes during the previous 30 days;]								
	(1) (4) or [(c) where there is no official restriction for health reasons and in and around which, in area of 50 km radius, there had case/outbreak of foot-and-mouth disease or rinderpest during the previous 90 days, and,								
		(d) where they have remained for at least 40 days before	e direct dispatch to the slaughterhouse	e;]					
	(¹) (⁸) or	[(d) where they have remained for at least 40 days be veterinary authority without coming into contact with a a slaughterhouse;]							
	II.2.4.	has been obtained from animals which:							
		(a) have been transported from their holdings in vehicles without contact with other animals which did not com							
		(b) at the slaughterhouse, have passed ante-mortem heal shown no evidence of the diseases referred to in poi		re slaughter and, in particular, have					
		(c) have been slaughtered on (dd/mm/yyyy)	or between (dd/mm/yyyy) and(dd/mm/yyyy) (⁶);					
	II.2.5.	has been obtained in an establishment around which, wir referred to in point II.2.1 during the previous 30 days or importation into the Union has been authorised only after and disinfection of the establishment under the control of	, in the event of a case/outbreak of di slaughter of all animals present, remove	sease, the preparation of meat for					
	II.2.6.								
	(1) either	[has been obtained and prepared without contact with o	ther meats not complying with the cor	nditions required in this certificate.]					
	(¹) (⁴) or	[contains [boneless meat] [and] [minced meat] (1), obta carcasses in which the main accessible lymphatic glar temperature above + 2 °C for at least 24 hours before th 6.0 when tested electronically in the middle of the longi	nds have been removed, which have be bones were removed and in which the	been submitted to maturation at a ne pH value of the meat was below					
		has been kept strictly separate from meat not conform production, de-boning and storage until it has been pac							
	(¹) (⁷) or	[contains [boneless meat] [and] [minced meat] (1), obta carcasses in which the main accessible lymphatic glar temperature above + 2 °C for at least 24 hours before to	nds have been removed, which have						
		has been kept strictly separate from meat not conform production, de-boning and storage until it has been pac							
II.3.	Animal	welfare attestation							
		dersigned official veterinarian, hereby certify, that the fresh ghterhouse before and at the time of slaughter or killing in							

▼<u>M1</u>

Date:

Stamp:

COUNTRY		Model OV				
II. Health information	II.a. Certificate reference number	II.b.				
Notes	1					
This certificate is meant for fresh meat, including minced meat, of do Fresh meat means all animal parts fit for human consumption whether f	,	nd caprine animals (Capra hircus).				
Part I:						
Box reference I.8: Provide the code of territory as appearing in Part	1 of Annex II to Regulation (EU) No 2	206/2010.				
Box reference I.11: Place of origin: name and address of the dispate	ch establishment.					
Box reference I.15: Registration number (railway wagons or containe case of unloading and reloading, the consignor must inform the BIP		or name (ship) is to be provided. In				
Box reference I.19: Use the appropriate HS code: 02.04, 02.06 or 05 column 5 "SG" of Part 1 of Annex II to Regulation (EU) No 206/2010						
Box reference I.20: Indicate total gross weight and total net weight.						
Box reference I.23: For containers or boxes, the container number a	nd the seal number (if applicable) sho	ould be included.				
 Box reference I.28: Nature of commodity: Indicate "carcass-whole", "carcass-side", "carcass-quarters", "cuts", "offal" or "minced meat". Minced meat is de-boned meat that has been minced into fragments and that must have been prepared exclusively from striated muscle (including the adjoining fatty tissues) except heart muscle. 						
Box reference I.28: Treatment type: If appropriate, indicate "de-bon freezing (mm/yy) of the cuts/pieces.	ed"; 'bone in"; "matured" and/or "mino	eed". If frozen, indicate the date of				
Part II:						
(¹) Keep as appropriate.						
(²) List of countries in the Annex to Decision 2007/453/EC.						
(3) Code of the territory as it appears in Part 1 of Annex II to Regulation	n (EU) No 206/2010.					
(4) Supplementary guarantees regarding meats from matured de-boned to Regulation (EU) No 206/2010, with the entry "A".	meat to be provided when required in o	column 5 "SG" of Part 1 of Annex II				
(5) Delete when the exporting country carries out vaccination against authorised to import into the Union matured de-boned meat which fu						
authorisation for importation into the Union of the third country, territor	6) Date or dates of slaughter. Imports of this meat shall not be allowed when obtained from animals slaughtered either prior to the date of authorisation for importation into the Union of the third country, territory or part thereof referred to in boxes I.7 and I.8, or during a period where restrictive measures have been adopted by the Union against imports of this meat from this third country, territory or part thereof.					
7) Supplementary guarantees regarding meats from matured de-boned meat to be provided when required in column 5 "SG" of Part 1 of Annex II to Regulation (EU) No 206/2010, with the entry "F". The matured de-boned meat shall not be authorised for importation into the Union until 21 days after the date of slaughter of the animals.						
(8) Alternative guarantee may be provided when allowed for by the entry "J" in column 5 "SG" of Part 1 of Annex II to Regulation (EU) No 206/2010.						
Official veterinarian						
Name (in capital letters):	Qualification and title	:				

Signature:'

		Mod	el POR				
	COUNTRY				Veterinary cer	tificate to EU	
	I.1. Consignor		I.2. Certific	ate reference nui	mber I.2.a.		
	Name		I.3. Central	Competent Auth	nority		
		Address			rity		
ent	Tel. No			competent Author			
gnn	I.5. Consignee		1.6.				
onsi	Name						
ed c	Address						
atch	Postal code						
lispa	Tel. No						
Part I: Details of dispatched consignment	I.7. Country ISO I.8. Region of origin code of or		I.9. Country destina			Code	
Det	I.11. Place of origin		I.12.				
rt :	Name Approval Address	number					
a a	Address						
	I.13. Place of loading		I.14. Date of	departure			
	I.15. Means of transport		I.16. Entry B	IP in EU			
	Aeroplane Ship	Railway wagon 🗌					
	Road vehicle Other						
	Identification:	I.17.					
	Documentary references:						
	I.18. Description of commodity			I.19. Commodi	ity code (HS code)		
				1.	.20. Quantity		
	I.21. Temperature of product			1.	.22. Number of package	es	
	Ambient Chile	ed 🗌	Frozen				
	I.23. Identification of container/seal number			I.	.24. Type of packaging		
	I.25. Commodities certified for: Human consumption						
,	1.26.		I.27. For imp	ort or admission	into EU		
	I.28. Identification of the commodities						
	(Scientific name) commodity type of packages weigh					Net weight	
		Abatto	ir Cutting p	olant Cold sto	re		

	COU	INTRY					Model	PO
	II.	Health	information		II.a. Certificate reference numbe	er	II.b.	
	II.1.	Public	: Health Attes	tation				
		(EC) N	lo 852/2004, (EC) No 853/		ereby certify th	ements of Regulations (EC) No 178/20 nat the meat of domestic swine describ	
		II.1.1			i] (¹) comes from (an) establishmer with Regulation (EC) No 852/2004		nting a programme based on the HAC	CF
II.1.1 the [meat] [minced meat] (¹) comes from (an) establishment(s) implementing a programme principles in accordance with Regulation (EC) No 852/2004; II.1.2 the meat has been obtained in compliance with the conditions set out in Section I of Annex No 853/2004; II.1.3 the meat fulfils the requirements of Regulation (EC) No 2075/2005 laying down specific rules							Section I of Annex III to Regulation (E	EC;
II.1.3 the meat fulfils the requirements of Regulation (EC) No 2075/2005 laying down specific rules on o Trichinella in meat, and in particular:								fo
		ethod with negative results]						
			(¹) or	[has been No 2075	,	ment in accord	dance with Annex II to Regulation (E	ΞC
			(¹) or	holding		een officially re	or fattening and slaughter, comes fron ecognized by the competent authority lation (EC) No 2075/2005;]	
		(¹) II.1.4			en produced in accordance with Se perature of not more than -18 °C;]	ection V of Anne	ex III to Regulation (EC) No 853/2004 a	an
		II.1.5		with Chapt			d post-mortem inspections carried out ection IV of Annex I to Regulation (E	
		II.1.6 (¹) either		cass or parts of the carcass have		d with a health mark in accordance w 154/2004;]	vit
			(¹) or		ekages of [meat] [minced meat] nce with Section I of Annex II to Re		marked with an identification mark No 853/2004;]	i
		II.1.7	the [meat] [n criteria for fo		(1) satisfies the relevant criteria set	out in Regulati	ion (EC) No 2073/2005 on microbiologi	ica
		II.1.8			live animals and products thereof and in particular Article 29, are fulfi		e residue plans submitted in accordar	nc
		II.1.9			t] (1) has been stored and transported of Annex III to Regulation (EC)		dance with the relevant requirements	; (
		(²) [II.1.10					Regulation (EC) No 853/2004 as regand Sweden of certain meat and eggs;]	rd
II.2. Animal Health attestation								
		I, the u	ındersigned of	ficial veterin	arian, hereby certify, that the fresh i	meat describe	d in Part I :	
		II.2.1	has been ob	tained in the	territory/ies with code:	(³) w	hich, at the date of issuing this certifica	ate
			(¹) either	- ' '	been free for 12 months from for sical swine fever, swine vesicular di		disease, rinderpest, African swine fev	ve

(1) or

[(a) (i) has been free for 12 months from rinderpest, African swine fever, [foot-and-mouth disease] ('), [classical swine fever] (') and [swine vesicular disease] ('), and

COUNTRY Model POR

II.	Health information			II.a. Certificate reference number	II.b.			
			1	has been considered free from [foot-and-mout] [swine vesicular disease] (¹), sincehad cases/outbreaks afterwards, and author Regulation (EC) No/, of	(dd/mm/yyyy) rised to export this meat (dd/mm/yyyy), a), without having by Commission and]		
			impo	orts of domestic animals vaccinated against tory;	these diseases are not p	permitted in this		
	11.2.2	has been obta	ained from a	animals that:				
		(¹) either		mained in the territory described under point II before slaughter;]	.2.1 since birth, or for at le	ast the last three		
		(¹) or	point II.2	een introduced on(dd/r 2.1, from the territory with code				
		(¹) or		een introduced on(dd/i 2.1, from the EU Member State(dd/i		described under		
	11.2.3	has been obta	ained from	animals coming from holdings:				
	point II.2.1,			ne animals present therein have been vacci	nated against the disease	es referred to in		
				, in an area of 10 km radius, there has been no e previous 40 days,	case/outbreak of the disea	ses referred to in		
		(c) that are r weeks;	ot subject	to prohibition as a result of an outbreak of p	porcine brucellosis during	the previous six		
	(1) (4)			g has been received that pigs are not fed with c he list established by the competent authority fo				
	11.2.4	has been obta	ained from	animals that:				
		(a) have rema	ained sepa	rate since birth from wild cloven-hoofed animal	s,			
			nouse with	orted from their holdings in vehicles, cleaned and disinfected before loading, to an appro thout contact with other animals which did not comply with the conditions set out in points II.3				
				e, have passed ante-mortem health inspection vn no evidence of the diseases referred to in po		slaughter and, in		
				red on(dd/mm/yyyy) or b (dd/mm/yyyy). (⁵);	etween	(dd/mm/yyyy)		
	II.2.5	of the diseas preparation of	es referred f meat for i	establishment around which, within a radius of to in point II.2.1 during the previous 40 days mportation into the Union has been authorised the total cleaning and disinfection of the establishment.	or, in the event of a case donly after slaughter of all	e of disease, the animals present,		
	II.2.6	has been obta certificate.	ained and p	repared without contact with other meats not c	omplying with the condition	ns required in this		
II.3.	Anima	ıl welfare attes	tation					
	been t			arian, hereby certify, that the fresh meat describ se before and at the time of slaughter or killing				

COUNTRY Model POR

II.	Health information	II.a. Certificate reference number	II.b.	

Notes

This certificate is meant for fresh meat, including minced meat, of domestic swine (Sus scrofa).

Fresh meat means all animal parts fit for human consumption whether fresh, chilled or frozen.

Part I:

- Box reference I.8: Provide the code of territory as appearing in Part 1 of Annex II to Regulation (EU) No 206/2010.
- Box reference I.11: Place of origin: name and address of the dispatch establishment.
- Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor must inform the BIP of entry into the Union.
- Box reference I.19: Use the appropriate HS code: 02.03, 02.06, 02.09, 05.04 or 15.01.
- Box reference I.20: Indicate total gross weight and total net weight.
- Box reference I.23: For containers or boxes, the container number and the seal number (if applicable) should be included.
- Box reference I.28: Nature of commodity: Indicate 'carcass-whole', 'carcass-side', 'carcass-quarters', 'cuts' or 'minced meat'.
 Minced meat is deboned meat that has been minced into fragments and that must have been prepared exclusively from striated muscle (including the adjoining fatty tissues) except heart muscle.
- Box reference I.28: Treatment type: If appropriate, indicate 'deboned'; 'bone in'; 'matured' and/or 'minced'. If frozen, indicate the date
 of freezing (mm/yy) of the cuts/pieces.

Part II:

- (1) Keep as appropriate.
- (2) Delete if the consignment is not intended for import into Finland or Sweden.
- (3) Code of the territory as it appears in Part 1 of Annex II to Regulation (EU) No 206/2010.
- (4) Supplementary guarantees to be provided when required in column 5 'SG' of Part 1 of Annex II to Regulation (EU) No 206/2010, with the entry 'D'.

Catering waste means: all waste from food intended for human consumption from restaurants, catering facilities or kitchens, including industrial kitchens and household kitchens of the farmer or persons tending pigs.

(°) Date or dates of slaughter. Imports of this meat shall not be allowed when obtained from animals slaughtered either prior to the date of authorisation for importation into the Union of the third country, territory or part thereof referred to in boxes I.7 and I.8, or during a period where restrictive measures have been adopted by the Union against imports of this meat from this third country, territory or part thereof.

Official	Official veterinarian									
	Name (in capital letters):	Qualification and title:								
	Date:	Signature:								
	Stamp:									

	Model EQU							
		Ountry			Veterinary certificate to EU			
	1.1.	Consignor	1.2. Certifica	ate reference numbe	er 1.2.a.			
		Name		I.3. Central	Competent Authorit	у		
		Address		I.4. Local C	ompetent Authority			
nen		Tel. No						
ign	1.5.	Consignee			I.6.			
suo:		Name						
ed		Address						
atch		Postal code						
disp		Tel. No						
Part I: Details of dispatched consignment	1.7.	Country ISO code	I.8. Region of origin	Code	I.9. Country destina		I.10. Region of Code destination	
Det	1.11	. Place of origin			l.12.			
ar I		Name Address	Approval number					
<u>a</u>		Address						
	I.13	3. Place of loading			I.14. Date of departure			
	1.15	. Means of transport			I.16. Entry BIP in EU			
		Aeroplane Sh	ip Railway wagon					
		Road vehicle Oth	er 🗌					
		Identification: Documentary references:		1.17.				
	I.18	3. Description of commodity				I.19. Commodity of	ode (HS code)	
						1.20.	Quantity	
	1.21	1. Temperature of product				1.22.	Number of packages	
		Ambient	Chiled		Frozen			
						1		
	1.23	3. Identification of container/s	eal number			1.24.	Type of packaging	
	1.25	5. Commodities certified for:						
		Human consumption						
	1.26	5.			I.27. For imp	ort or admission into	EU	
I.28. Identification of the commodities								
	(Nature of Approximately	oroval nu	umber establis	hments	Number Net of packages weight	
			Abatto	oir C	Cutting plant	Cold store		

COUNTRY Model EQU II. Health information II.a. Certificate reference number II.b. II.1. **Public Health Attestation** I, the undersigned official veterinarian, declare that I am aware of the relevant requirements of Regulations (EC) No 178/2002, (EC) No 852/2004, (EC) No 853/2004 and (EC) No 854/2004 and hereby certify that the meat of domestic solipeds described in Part I was produced in accordance with those requirements, in particular that: Part II: Certification the meat comes from (an) establishment(s) implementing a programme based on the HACCP principles in accordance with Regulation (EC) No 852/2004; 11.1.1 the meat has been obtained in compliance with the conditions set out in Section I of Annex III to Regulation (EC) II.1.2 No 853/2004: the meat fulfils the requirements of Regulation (EC) No 2075/2005 laying down specific rules on official controls II.1.3 for Trichinella in meat, and in particular, has been subject to an examination by a digestion method with negative the meat has been found fit for human consumption following ante and post-mortem inspections carried out in accordance with Chapter II of Section I and Chapters III and IX of Section IV of Annex I to Regulation (EC) No 854/2004; II.1.5 (1) either [the carcass or parts of the carcass have been marked with a health mark in accordance with Chapter III of Section I of Annex I to Regulation (EC) No 854/2004;] (1) or [the packages of meat have been marked with an identification mark in accordance with Section I of Annex II to Regulation (EC) No 853/2004;] II.1.6 the meat satisfies the relevant criteria set out in Regulation (EC) No 2073/2005 on microbiological criteria for foodstuffs: II.1.7 the guarantees covering live animals and products thereof provided by the residue plans submitted in accordance with Directive 96/23/EC, and in particular Article 29 thereof, are fulfilled; the meat has been stored and transported in accordance with the relevant requirements of Section I of Annex III to II.1.8 Regulation (EC) No 853/2004. II.2. **Animal Health attestation** I, the undersigned official veterinarian, hereby certify, that the fresh meat described in Part I: II.2.2 has been obtained from domestic solipeds, which: (1) either [have remained in the territory described under point II.2.1 since birth, or for at least the last three months before slaughter;] [have been introduced on (1) or(dd/mm/yyyy) into the territory described under point II.2.1, from the territory with code: (2) that at that date was authorised to export this fresh meat to the Union;] (1) or II.2.3 has been obtained from animals which were slaughtered on (dd/mm/yyyy) or between (dd/mm/yyyy) and (dd/mm/yyyy) (3) in a slaughterhouse around which, within a radius of 10 km, there has been no case/outbreak of African horse sickness or glanders during the previous 40 days or, in the event of a case of such diseases, the preparation of meat for importation into the Union has been authorised only after slaughter of all animals present, removal of all meat, and the total cleaning and disinfection of the establishment under the control of an official veterinarian;

COUNTRY Model EQU

II.	Health information	II.a. Certificate reference number	II.b.

II.2.4 has been obtained and prepared without contact with other meats not complying with the conditions required in this certificate.

II.3. Animal welfare attestation

I, the undersigned official veterinarian, hereby certify that the fresh meat described in this certificate derives from animals which have been treated in the slaughterhouse before and at the time of slaughter or killing in accordance with the relevant provisions of Union legislation.

Notes

This certificate is meant for fresh meat, excluding minced meat, of domestic solipeds (Equus caballus, Equus asinus and their cross-breeds).

Fresh meat means all animal parts fit for human consumption whether fresh, chilled or frozen.

Part I:

- Box reference I.8: Provide the code of territory as appearing in Part 1 of Annex II to Regulation (EU) No 206/2010.
- Box reference I.11: Place of origin: name and address of the dispatch establishment.
- Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor must inform the BIP of entry into the Union.
- Box reference I.19: Use the appropriate HS code: 02.05, 02.06 or 05.04.
- Box reference I.20: Indicate total gross weight and total net weight.
- Box reference I.23: For containers or boxes, the container number and the seal number (if applicable) should be included.
- Box reference I.28: Nature of commodity: Indicate 'carcass-whole', 'carcass-side', 'carcass-quarters' or 'cuts'.
- Box reference I.28: Treatment type: If appropriate, indicate 'deboned'; 'bone in' and/or 'matured'. If frozen, indicate the date of freezing (mm/yy) of the cuts/pieces.

Part II:

- (1) Keep as appropriate.
- (2) Code of the territory as it appears in Part 1 of Annex II to Regulation (EU) No 206/2010.
- (3) Dates: imports of this meat shall not be authorised when obtained from animals slaughtered either prior to the date of authorisation for importation into the Union of the third country, territory or part thereof referred to in boxes I.7 and I.8, or during a period where restrictive measures have been adopted by the Union against imports of this meat from this third country, territory or part thereof.

Official v	Official veterinarian					
	Name (in capital letters):	Qualification and title:				
	Date:	Signature:				
	Stamp:					

		lel RUF				
	COUNTRY	Veterinary certificate to EU I.2. Certificate reference number I.2.a.				
	I.1. Consignor	1.2. Certificate reference flumber 1.2.a.				
	Name	I.3. Central Competent Authority				
	Address	I.4. Local Competent Authority				
nent	Tel. No	· · · · · · · · · · · · · · · · · · ·				
ignn	I.5. Consignee	1.6.				
ons	Name					
o pa	Address					
atch	Postal code					
lispa	Tel. No					
Part I: Details of dispatched consignment	I.7. Country ISO I.8. Region Code of origin code of origin	I.9. Country of ISO I.10. Region of Code destination code				
Det	I.11. Place of origin	I.12.				
벌	Name Approval number					
<u>%</u>	Address					
	I.13. Place of loading	I.14. Date of departure				
	I.15. Means of transport	I.16. Entry BIP in EU				
	Aeroplane Ship Railway wagon					
	Road vehicle Other					
	Identification:	1.17.				
	Documentary references:					
	I.18. Description of commodity	I.19. Commodity code (HS code)				
		I.20. Quantity				
	I.21. Temperature of product	I.22. Number of packages				
	Ambient ☐ Chiled ☐	Frozen				
		_				
	I.23. Identification of container/seal number	I.24. Type of packaging				
	I.25. Commodities certified for:					
	Human consumption					
	1.26.	I.27. For import or admission into EU				
	I.28. Identification of the commodities					
	(Scientific name) commodity type	oroval number establishments Number Net of packages weight				
	Abatto	oir Cutting plant Cold store				

COUNTRY Model RUF II. Health information II.a. Certificate reference number II.b. II.1. **Public Health Attestation** I, the undersigned official veterinarian, declare that I am aware of the relevant requirements of Regulations (EC) No 178/2002, (EC) No 852/2004, (EC) No 853/2004, (EC) No 854/2004 and (EC) No 999/2001 and hereby certify that the meat of farmed animals of the order Artiodactyla (excluding bovine animals (including *Bison* and *Bubalus* species and their cross-breeds), *Ovis aries, Capra hircus*, Suidae and Tayassuidae), and of the families Rhinocerotidae and Part II: Certification Elephantidae described in Part I was produced in accordance with those requirements, in particular that: the meat comes from (an) establishment(s) implementing a programme based on the HACCP principles in accordance with Regulation (EC) No 852/2004; II.1.2 the meat has been obtained in accordance with the conditions set out in Section III of Annex III to Regulation (EC) No 853/2004: II.1.3 the meat has been found fit for human consumption following ante and post-mortem inspections carried out in accordance with Chapter II of Section I and Chapters VII and IX of Section IV of Annex I to Regulation (EC) No 854/2004; II.1.4 (1) either [the carcass or parts of the carcass have been marked with a health mark in accordance with Chapter III of Section I of Annex I to Regulation (EC) No 854/2004;] (1) or [the packages of meat have been marked with an identification mark in accordance with Section I of Annex II to Regulation (EC) No 853/2004;] the meat satisfies the relevant criteria set out in Regulation (EC) No 2073/2005 on microbiological criteria for foodstuffs; II.1.6 the guarantees covering live animals and products thereof provided by the residue plans submitted in accordance with directive 96/23/EC, and in particular Article 29 thereof, are fulfilled (1) (2) [II.1.7 with regard to Chronic Wasting Disease (CWD): This product contains or is derived exclusively from meat, excluding offal and spinal cord, of farmed cervid animals which have been examined for Chronic Wasting Disease by histopathology, immunohistochemistry or other diagnostic method recognised by the competent authority with negative results and is not derived from animals coming from a herd where Chronic Wasting Disease has been confirmed or is officially suspected.] the meat has been stored and transported in accordance with the relevant requirements of Section I of Annex III to Regulation (EC) No 853/2004.

II.2. Animal Health attestation

I, the undersigned official veterinarian, hereby certify, that the fresh meat described in Part I:

- - (a) has been free for 12 months from rinderpest, and during the same period no vaccination against this disease has taken place, and
- (') either [(b) has been free for 12 months from foot-and-mouth disease, and during the same period no vaccination against this disease has taken place;]
- (¹) (⁴) or [(b) vaccination programmes against foot-and-mouth disease are being officially carried out and controlled in domestic bovine animals;]

COUNTRY Model RUF

II.	II. Health information		II.a. Certificate reference numb	rtificate reference number II.b.		
	II.2.2	has been obtained	from animals that:			
			ave remained in the territory described uponths before slaughter;]	under point l	I.2.1 since birth, or for at least the last three	
		(¹) or [have been introduced on				
	II.2.3	has been obtained	from animals coming from holdings:			
		(a) in which none of the animals present therein have been vaccinated against [foot-and-mouth d or] (5) rinderpest,				
					seases transmissible to humans or animals utbreak of brucellosis during the previous six	
	(¹) either		which in an area of 10 km radius, there hing the previous 30 days,]	nas been no	case/outbreak of foot-and-mouth disease or	
	(¹) (⁴) or [(c) where there is no official restriction for health reasons and in and around which in an area of 50 km radius has been no case/outbreak of foot-and-mouth disease or rinderpest during the previous 90 days, and					
		(d) where the anim	nals have remained for at least 40 days b	oefore direct	dispatch to the slaughterhouse;]	
	II.2.4	has been obtained	from animals:			
	(¹) either				aned and disinfected before loading, to an did not comply with the conditions mentioned	
			aughterhouse, have passed ante-morter lar, have shown no evidence of the disea		pection during the 24 hours before slaughter to in point II.2.1, and	
			een slaughtered on(dd/mm/yyyy		n/yyyy) or between	
	(¹) or		een slaughtered on the holding of ori r the holding, who has provided a writte		g authorisation by an official veterinarian that:	
			ion an unacceptable risk would have bee isport of the animals to an slaughterhous		he welfare of the animals or to their handlers	
		 the holdin animals, 	g had been inspected and authorised	by the com	petent authority for the slaughter of game	
			is have passed the ante-mortem health i ar, have shown no evidence of the diseas		uring the 24 hours before the slaughter and, to in point II.2.1,	
		the anima (dd/mm/y)			(dd/mm/yyyy) and	
		the bleedi	ng of the animals was performed correct	ly, and		
		the slaugh	tered animals were eviscerated within th	ree hours of	the time of slaughter, and	
	(b) the carcasses of which have been transported to the approved slaughterhouse under hygienic conditions a where more than one hour elapsed since the time of slaughter, a temperature of between 0 °C and + 4 °C been found on the arrival of the vehicle used for the transport;]					
	(¹) (²) II.2.5	[has been obtaine hoofed animals;]	d from animals that have remained since	birth or for t	he last 3 months separate from wild cloven-	

COUNTRY Model RUF

II.	Health information			II.a. Certificate reference number	II.b.		
	of the diseases referred preparation of meat for in		s referred meat for in	establishment around which, within a radius to in point II.2.1 during the previous 30 days mportation into the Union has been authorised I the total cleaning and disinfection of the es	s or, in the event of a case of disease, the d only after slaughter of all animals present,		
	II.2.7						
		(¹) either	[has bee required	n obtained and prepared without contact with other meats not complying with the conditions above.]			
		(¹) (⁴) or	carcasse submitte removed	ins boneless meat, obtained only from de-boned meat other than offal that was obtained from sees in which the main accessible lymphatic glands have been removed, which have been ted to maturation at a temperature above $+2^{\circ}\mathrm{C}$ for at least 24 hours before the bones were ed and in which the pH value of the meat was below 6.0 when tested electronically in the e of the longissimus-dorsi muscle after maturation and before de-boning, and			
			certificat	has been kept strictly separate from meat not conforming to the requirements set out in t certificate during all stages of its production, de-boning and storage until it has been packed boxes or cartons for further storage in dedicated areas.]			
		(¹) (⁸) or	carcasse	s boneless meat, obtained only from de-boned es in which the main accessible lymphatic gla d to maturation at a temperature above + 2 °C I, and	ands have been removed, which have been		
			certificat	n kept strictly separate from meat not confo e during all stages of its production, de-boni cartons for further storage in dedicated areas.	ng and storage until it has been packed in		

Notes

This certificate is meant for fresh meat, excluding offal and minced meat, of wild animals of the order Artiodactyla (excluding bovine animals (including *Bison* and *Bubalus* species and their cross-breeds), *Ovis aries, Capra hircus*, Suidae and Tayassuidae), and of the families Rhinocerotidae and Elephantidae, that are domestically kept or bred since birth or for the last three months in farms.

Fresh meat means all animal parts fit for human consumption whether fresh, chilled or frozen.

Part I:

- Box reference I.8: Provide the code of territory as appearing in Part 1 of Annex II to Regulation (EU) No 206/2010.
- Box reference I.11: Place of origin: name and address of the dispatch establishment.
- Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor must inform the BIP of entry into the Union.
- Box reference I.19: Use the appropriate HS code: 02.06, 02.08.90 or 05.04.
- Box reference I.20: Indicate total gross weight and total net weight.
- Box reference I.23: For containers or boxes, the container number and the seal number (if applicable) should be included.
- Box reference I.28: Nature of commodity: Indicate 'carcass-whole', 'carcass-side', 'carcass-quarters', or 'cuts'.
- Box reference I.28: Treatment type: If appropriate, indicate 'deboned'; 'bone in' and/or 'matured'. If frozen, indicate the date of freezing (mm/yy) of the cuts/pieces.

CC	COUNTRY Model RUI						
11.	Health information	II.a. Certificate reference number	II.b.				
Pa	rt II:						
(1)	Keep as appropriate.						
٠,,			provided when required in column 5 'SG' of Part				
(3)	Code of the territory as it appears in Par	t 1 of Annex II to Regulation (EU) No 206	5/2010.				
	Part 1 of Annex II to Regulation (EU) N	lo 206/2010 with the entry 'A'.	be provided when required in column 5 'SG' of				
(5)	(5) Delete when the exporting country carries out vaccination against foot-and-mouth disease with serotypes A, O or C, and this country is allowed for import into the Union matured de-boned meat which fulfils the supplementary guarantees described under footnote (4).						
(⁶)	(°) Date or dates of slaughter. Imports of this meat shall not be authorised when obtained from animals slaughtered either prior to the date of authorisation for importation into the Union of the third country, territory or part thereof referred to in boxes I.7 and I.8, or during a period where restrictive measures have been adopted by the Union against imports of this meat from this third country, territory or part thereof.						
(⁷)	Not necessary for farmed game animals	s kept permanently in Arctic regions.					
(8)		010, with the entry 'F'. The matured de-b	provided when required in column 5 'SG' of Part 1 oned meat shall not be authorised for importation				
Off	icial veterinarian						
	Name (in capital letters):	Qualific	cation and title:				
	Date:	Signati	ure:				
	Stamp:						

				Mode	I RUW				
		UNTRY			Veterinary certificate to EU				
	I.1.	Consignor			I.2. Certificate reference number I.2.a.				
		Name			I.3. Central Competent Authority				
		Address			I.4. Local C	ompetent Author	itv		
Jent		Tel. No							
ligi	I.5.	Consignee			I.6.				
onsi		Name							
o pa		Address							
atch	Postal code								
lisp		Tel. No							
Part I: Details of dispatched consignment	1.7.	Country ISO code	I.8. Region Control of origin	Code	I.9. Country destina			.10. Region of destination	Code
Det	l.11.	. Place of origin			l.12.				
rt	Name Approval number								
<u>a</u>	Address								
	I.13	I.13. Place of loading			I.14. Date of departure				
	I.15. Means of transport				I.16. Entry BIP in EU				
	Aeroplane Ship Railway wagon Road vehicle Other								
		Identification: Documentary references:			l.17.				
	I.18	. Description of commodity				I.19. Commodit	y cod	e (HS code)	
					ļ	1.:	20. Qu	uantity	
	1.21	. Temperature of product				1.3	22. Nu	ımber of package	s
		Ambient	Chiled		Frozen				
	1.23	3. Identification of container/se	eal number			1.3	24. Ty _l	pe of packaging	
	1.25	5. Commodities certified for:						·	
		Human consumption							
	1.26.			I.27. For import or admission into EU					
	I.28. Identification of the commodities								
	Species Nature of Treatment App (Scientific name) commodity type					establishments		Number of packages	Net weight
	Abattoir Cutting plant Cold store								

(1) either

this disease has taken place;]

COUNTRY Model RUW II. Health information II.a. Certificate reference number II.b. II.1. **Public Health Attestation** I, the undersigned official veterinarian, declare that I am aware of the relevant requirements of Regulations (EC) No 178/2002, (EC) No 852/2004, (EC) No 853/2004 and (EC) No 854/2004 and hereby certify that the fresh meat of wild animals of the order Artiodactyla (excluding bovine animals (including *Bison* and *Bubalus* species and their cross-breeds), *Ovis aries, Capra hircus,* Suidae and Tayassuidae), and of the families Rhinocerotidae and Elephantidae described in Part II: Certification Part I was produced in accordance with those requirements, in particular that: the meat comes from (an) establishment(s) implementing a programme based on the HACCP principles in accordance with Regulation (EC) No 852/2004; II.1.2 the meat has been obtained in compliance with the conditions set out in Section IV of Annex III to Regulation 853/2004, and in particular: (i) before skinning, it has been stored and handled separately from other food and not frozen; and (ii) after skinning, it has undergone a final inspection as referred to in point II.1.4; (1) II.1.3 [in the case of susceptible species, the meat fulfils the requirements of Regulation (EC) No 2075/2005 laying down specific rules on official controls for Trichinella in meat:1 the meat has been found fit for human consumption following a post-mortem inspection carried out in accordance with Chapter II of Section I and Chapters VIII and IX of Section IV of Annex I to Regulation (EC) No 854/2004; II.1.4 [in the case of large wild game, the carcass or parts of the carcass have been marked with a health II.1.5 (1) either mark in accordance with Chapter III of Section I of Annex I to Regulation (EC) No 854/2004;] [the packages of meat have been marked with an identification mark in accordance with Section I of (1) or Annex II to Regulation (EC) No 853/2004;] the meat satisfies the relevant criteria set out in Regulation (EC) No 2073/2005 on microbiological criteria for II.1.6 foodstuffs; the guarantees covering live animals and products thereof provided by the residue plans submitted in accordance II.1.7 with Directive 96/23/EC, and in particular Article 29 thereof, are fulfilled. (1) (2) [II.1.8 with regard to Chronic Wasting Disease (CWD): This product contains or is derived exclusively from meat, excluding offal and spinal cord, of wild cervid animals which have been examined for Chronic Wasting Disease by histopathology, immunohistochemistry or other diagnostic method recognised by the competent authority with negative results and is not derived from animals coming from a region where Chronic Wasting Disease has been confirmed in the last three years or is officially suspected.] the meat has been stored and transported in accordance with the relevant requirements of Section I of Annex III to Regulation (EC) No 853/2004. 11.2. **Animal Health attestation** I, the undersigned official veterinarian, hereby certify, that the fresh meat described in Part I: II.2.1 has been obtained in the territory/ies with code:(3) which, at the date of issuing this certificate: (a) has been free for 12 months from rinderpest, and during the same period no vaccination against this disease has taken place, and

[(b) has been free for 12 months from foot-and-mouth disease, and during the same period no vaccination against

COUNTRY Model RUW

II. Health information		II.a. C	ertificate reference number	II.b.			
(¹) or	having h	ad cases/outbreaks		(dd/mm/yyyy), without these animals by Commission Regulation			
(¹) (⁴) or		on programmes ag bovine animals;]	gainst foot-and-mouth disease are b	eing officially carried out and controlled in			
II.				(dd/mm/yyyy) and to in point II.2.1, and the killing took place:			
	, ,		needs 20 km from the borders of a country or part thereof, which is not authorised during this this fresh meat into the Union,				
	(b) in an are point II.2.		e last 60 days, there has been no	restrictions for the diseases referred to in			
II	game-handlin diseases refe of meat for im	g establishment ar red to in point II.2. portation into the U	ound which, within a radius of 10 kill during the previous 30 days or, in the	s soon as possible for chilling to an approved n, there has been no case/outbreak of the e event of a case of disease, the preparation moval of all meat, and the total cleaning and arian;			
Ш	.2.4						
	(¹) either	[has been obtain required above.]	ed and prepared without contact with o	other meats not complying with the conditions			
	(¹) (⁴) or	carcasses in wh submitted to ma removed and in	ich the main accessible lymphatic gla turation at a temperature above +2°C	meat other than offal that was obtained from ands have been removed, which have been for at least 24 hours before the bones were below 6.0 when tested electronically in the n and before de-boning, and			
		certificate during		orming to the requirements set out in this ing and storage until it has been packed in i.]			
	(¹) (⁶) or	carcasses in wh	ich the main accessible lymphatic gla	meat other than offal that was obtained from ands have been removed, which have been for at least 24 hours before the bones were			
		certificate during		orming to the requirements set out in this ing and storage until it has been packed in]			
Notes							
	ata is meant for freezh	n meat evoluting o	offal and minced meat of wild animals of the state of the	s of the order Artiodectule (excluding boying			

This certificate is meant for fresh meat, excluding offal and minced meat, of wild animals of the order Artiodactyla (excluding bovine animals (including *Bison* and *Bubalus* species and their cross-breeds), *Ovis aries, Capra hircus*, Suidae and Tayassuidae), and of the families Rhinocerotidae and Elephantidae that are killed or hunted in the wild.

Fresh meat means all animal parts fit for human consumption whether fresh, chilled or frozen.

 $After importation, unskinned carcasses \ must be \ conveyed \ without \ delay \ to \ the \ processing \ establishment \ of \ destination.$

COUNTRY	Model RUW
COUNTRY	Wodel how

II.	Health information	II.a. Certificate reference number	II.b.

Part I:

- Box reference I.8: Provide the code of territory as appearing in Part 1 of Annex II to Regulation (EU) No 206/2010.
- Box reference I.11: Place of origin: name and address of the dispatch establishment.
- Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor must inform the BIP of entry into the Union.
- Box reference I.19: Use the appropriate HS code: 02.01, 02.02, 02.04, 02.06, 02.08.90 or 05.04.
- Box reference I.20: Indicate total gross weight and total net weight.
- Box reference I.23: For containers or boxes, the container number and the seal number (if applicable) should be included.
- Box reference I.28: Nature of commodity: Indicate 'carcass-whole', 'carcass-side', 'carcass-quarters' or 'cuts'.
- Box reference I.28: Treatment type: If appropriate, indicate 'matured' or 'unskinned'. If frozen, indicate the date of freezing (mm/yy) of the cuts/pieces.
- Box reference I.28: Abattoir: any abattoir or game handling establishment.

Part II:

- (1) Keep as appropriate
- (2) Supplementary guarantees regarding fresh meat obtained from cervids to be provided when required in column 5 'SG' of Part 1 of Annex II to Regulation (EU) No 206/2010, with the entry 'G'.
- (3) Code of the territory as it appears in Part 1 of Annex II to Regulation (EU) No 206/2010.
- (4) Supplementary guarantees regarding meat from matured de-boned meat to be provided when required in column 5 'SG' of Part 1 of Annex II to Regulation (EU) No 206/2010 with the entry 'A'.

The matured de-boned meat shall not be authorised for importation into the Union until 21 days after the date of killing of the animals

- (5) Dates. Imports of this meat shall not be authorised when obtained from animals killed or hunted either prior to the date of authorisation for importation into the Union of the third country, territory or part thereof referred to in boxes I.7 and I.8, or during a period where restrictive measures have been adopted by the Union against imports of this meat from this third country, territory or part thereof.
- (6) Supplementary guarantees regarding meats from matured de-boned meat to be provided when required in column 5 'SG' of Part 1 of Annex II to Regulation (EU) No 206/2010, with the entry 'F'. The matured de-boned meat shall not be allowed for importation into the Union until 21 days after the date of slaughter of the animals.

Official veterinarian		
Name (in capital letters):	Qualification and title:	
Date:	Signature:	
Stamp:		

	Model SUF							
		DUNTRY			10 0	-1	er 1.2.a.	
	1.1.	Consignor			I.2. Certificate reference number I.2.a.			
		Name			I.3. Central	Competent Author	ty	
l		Address			I.4. Local C	competent Authority	,	
nent		Tel. No						
ignr	1.5.	Consignee			I.6.			
suo:		Name						
ed		Address						
atch		Postal code						
disp		Tel. No						
Part I: Details of dispatched consignment	1.7.	Country ISO of origin code	I.8. Region Co of origin	de	I.9. Country destina		I.10. Region of Code destination	
Det	1.11	. Place of origin			1.12.			
art	Name Approval number							
۵	Address							
	1.13	I.13. Place of loading			I.14. Date of departure			
	I.15. Means of transport				I.16. Entry BIP in EU			
	Aeroplane Ship Railway wagon							
	Road vehicle Other							
		Identification: Documentary references:			1.17.			
	140						- 1- (110 1.)	
	1.10	Description of commodity				I.19. Commodity	code (HS code)	
						1.20	. Quantity	
	1.21	1. Temperature of product				1.22	. Number of packages	
		Ambient	Chiled		Frozen			
	1.23	3. Identification of container/se	eal number			1.24	. Type of packaging	
	1.25	5. Commodities certified for:				<u> </u>		
		Human consumption						
	1.26.				I.27. For import or admission into EU			
	I.28. Identification of the commodities							
	(Species Nature Scientific name) commo	dity type			establishments	Number Net of packages weight	
			Al	oattoi	r Cutting p	olant Cold store		

COUNTRY Model SUF II. Health information II.a. Certificate reference number II.b. II.1. **Public Health Attestation** I, the undersigned official veterinarian declare that I am aware of the relevant provisions of Regulations (EC) No 178/2002, (EC) No 852/2004, (EC) No 853/2004 and (EC) No 854/2004 and hereby certify that the meat of farmed non-domestic animals belonging to the Suidae, Tayassuidae, or Tapiridae families described in Part I was produced in accordance with those requirements, in particular that: Part II: Certification the meat comes from (an) establishment(s) implementing a programme based on the HACCP principles in accordance with Regulation (EC) No 852/2004; the meat has been obtained in compliance with the conditions set out in Section III of Annex III to Regulation (EC) the meat fulfils the requirements of Regulation (EC) No 2075/2005 laying down specific rules on official controls II.1.3 for Trichinella in meat, and in particular, has been subject to an examination by a digestion method with negative the meat has been found fit for human consumption following ante and post-mortem inspections carried out in accordance with, Chapter II of Section I and, Chapters VII and IX of Section IV of Annex I to Regulation (EC) No 854/2004; Ithe carcass or parts of the carcass have been marked with a health mark in accordance with II.1.5 (1) either Chapter III of Section I, of Annex I to Regulation (EC) No 854/2004;] (1) or [the packages of meat have been marked with an identification mark in accordance with Section I of Annex II to Regulation (EC) No 853/2004;] the meat satisfies the relevant criteria set out in Regulation (EC) No 2073/2005 on microbiological criteria for II.1.6 foodstuffs: 11.1.7 the guarantees covering live animals and products thereof provided by the residue plans submitted in accordance with Directive 96/23/EC, and in particular Article 29 thereof, are fulfilled; II.1.8 the meat has been stored and transported in accordance with the relevant requirements of Section I of Annex III to Regulation (EC) No 853/2004. 11.2. **Animal Health attestation** I, the undersigned official veterinarian, hereby certify, that the fresh meat described in Part I: (1) either [(a) has been free for 12 months from foot-and-mouth disease, rinderpest, African swine fever, classical swine fever, swine vesicular disease, and] (1) or [(a) (i) has been free for 12 months from rinderpest, African swine fever, [foot-and-mouth disease] (¹), [classical swine fever] (1) and [swine vesicular disease] (1), and (ii) has been considered free from [foot-and-mouth disease] (1), [classical swine fever] (1) and [swine vesicular disease] (1), since (dd/mm/yyyy), without having had cases/outbreaks afterwards, and authorised to export this meat by Commission (b) during the last 12 months no vaccination against these diseases have been carried out and imports of domestic animals vaccinated against these diseases are not permitted in this II.2.2 has been obtained from animals that:

[have remained in the territory described under point II.2.1 since birth, or for at least the last three

(1) either

months before slaughter;]

COUNTRY Model SUF

II.	Health	Health information		II.a. Certificate reference number	II.b.		
		(¹) or	point II.2	een introduced on			
	11.2.3	has been obtair	nas been obtained from animals coming from holdings:				
		(a) in which no point II.2.1,	 (a) in which none of the animals present therein have been vaccinated against the diseases referred to in point II.2.1, 				
				in an area of 10 km radius, there has been no e previous 40 days,	case/outbreak of the diseases referred to in		
			holdings	rinary inspections are carried out to diagnose d are not subject to prohibition as a result of an			
	11.2.4	has been obtair	ned from	animals which:			
		(¹) either	to a	e been transported from their holdings in vehicl n approved slaughterhouse without contact with ditions mentioned above,			
				ne slaughterhouse, have passed ante-mortem highter and, in particular, have shown no eviden			
				e been slaughtered on(dd. /mm/yyyy) and(dd/mm/			
		(¹) or		e been slaughtered on the holding of origin, follo consible for the holding, who has provided a writ			
			_	in his opinion an unacceptable risk would have to their handlers by the transport of the animals			
			_	the holding had been inspected and authorised of game,	by the competent authority for the slaughter		
			_	the animals have passed the ante-mortem he the slaughter and, in particular, have shown point II.2.1,			
			-	the animals were slaughtered between (dd/mm/yyyy), (³)	(dd/mm/yyyy) and		
			_	the bleeding of the animals was performed corr	rectly, and		
			_	the slaughtered animals were eviscerated within	in three hours of the time of slaughter, and		
			con	r carcasses have been transported to the a ditions and, where more than one hour of perature of between 0 °C and + 4 °C has been the transport;]	elapsed since the time of slaughter, a		
	II.2.5	has been obtair	ned from	animals that have remained separate since birt	th from wild cloven-hoofed animals;		
	II.2.6	of the diseases preparation of r	referred neat for	n establishment around which, within a radius of to in point II.2.1 during the previous 40 days importation into the Union has been authorised the total cleaning and disinfection of the establishment.	or, in the event of a case of disease, the donly after slaughter of all animals present,		
	II.2.7	has been obtair certificate.	ed and p	orepared without contact with other meats not co	emplying with the requirements set out in this		

COUNTRY	Model SUF

II.	Health information	II.a. Certificate reference number	II.b.					
II.3.	Animal welfare attestation							
	I, the undersigned official veterinarian, hereby certify, that the fresh meat described in Part I derives from animals which have been treated in the slaughterhouse before and at the time of slaughter or killing in accordance with the relevant provisions							

Notes

This certificate is meant for fresh meat, excluding offal and minced meat, of wild animals belonging to the Suidae, Tayassuidae, or Tapiridae families that are domestically kept or bred since birth in farms.

Fresh meat means all animal parts fit for human consumption, whether fresh, chilled or frozen.

Part I:

- Box reference I.8: Provide the code of territory as appearing in Part 1 of Annex II to Regulation (EU) No 206/2010.
- Box reference I.11: Place of origin: name and address of the dispatch establishment.
- Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor must inform the BIP of entry into the Union.
- Box reference I.19: Use the appropriate HS code: 02.03, 02.08.90 or 05.04.
- Box reference I.20: Indicate total gross weight and total net weight.
- Box reference I.23: For containers or boxes, the container number and the seal number (if applicable) should be included.
- Box reference I.28: Nature of commodity: Indicate 'carcass-whole', 'carcass-side', 'carcass-quarters' or 'cuts'.
- Box reference I.28: Treatment type: If appropriate indicate deboned, or bone-in. If frozen, indicate the date of freezing (mm/yy) of the cuts/pieces.

Part II:

- (1) Keep as appropriate
- (2) Code of the territory as it appears in Part 1 of Annex II to Regulation (EU) No 206/2010.
- (3) Date or dates of slaughter. Imports of this meat shall not be allowed when obtained from animals slaughtered either prior to the date of authorisation for importation into the Union of the third country, territory or part thereof referred to in boxes I.7 and I.8, or during a period where restrictive measures have been adopted by the Union against imports of this meat from this third country, territory or part thereof.

Official veterinarian	Official veterinarian						
Name (in	capital letters):		Qualification and title:				
Date:			Signature:				
Stamp:							

	Model SUW									
		Consigner				LO Cortifio	ate reference num		Veterinary certif	icate to Et
	1.1.	Consignor Name				i.z. Certific	ate reference num	iber	.2.a.	
		Address				I.3. Central	Competent Author	rity		
_						I.4. Local C	Competent Authori	ty		
nen						1.6.				
ign	1.5.	-				1.0.				
čouš		Name								
ped		Address Postal code								
atcl		Tel. No								
disk	17	Country	ISO	I.O. Danian	Code	10 Causatus	y of ISO	110	. Decien of	Code
Part I: Details of dispatched consignment	1.7.	of origin	code	I.8. Region of origin	Code	I.9. Country destina		1.10	D. Region of destination	Code
Det	1.11.	. Place of origin				I.12.				
art		Name Address		Approval number						
<u> </u>		Address								
	1.13	. Place of loading			I.14. Date of departure					
	1.15	. Means of transpor	rt			I.16. Entry B	IP in EU			
		Aeroplane Ship Railway wagon								
		Road vehicle								
		Identification:				I.17.				
		Documentary refe	rences:							
	I.18	. Description of con	nmodity				I.19. Commodity	y code ((HS code)	
							1.2	20. Quai	ntity	
	1.21	. Temperature of pr	oduct				1.2	22. Num	ber of packages	
		Ambient		Chiled		Frozen	1			
	1.23	. Identification of co	ontainer/s	eal number			1.2	24. Type	of packaging	
	1.25	. Commodities cert	ified for:							
		Human consumpt	ion 🗌							
	1.26. I.27. For it					I.27. For imp	port or admission i	nto EU		
	1.28	. Identification of th	e commo	dities						
	(5	Species Scientific name)	Nature commo		Арр	roval number e	establishments		Number packages	Net weight
					Abatto	ir Cutting p	olant Cold store	9		

COUNTRY Model SUW II. Health information II.a. Certificate reference number II.b. II.1. **Public Health Attestation** I, the undersigned official veterinarian declare that I am aware of the relevant requirements of Regulations (EC) No 178/2002, (EC) No 852/2004,(EC) No 853/2004 and (EC) No 854/2004 and hereby certify that the meat of wild animals belonging to the Suidae, Tayassuidae, or Tapiridae families described in Part I was produced in accordance with those requirements, in particular that Part II: Certification the meat comes from (an) establishment(s) implementing a programme based on the HACCP principles in accordance with Regulation (EC) No 852/2004; II.1.2 the meat has been obtained in accordance with Section IV of Annex III to Regulation (EC) No 853/2004, an in particular (i) before skinning, it has been stored and handled separately from other food and not frozen; and (ii) after skinning, it has undergone a final inspection as referred to in point II.1.4; the meat fulfils the requirements of Regulation (EC) No 2075/2005 laying down specific rules on official controls for Trichinella in meat, and in particular, has been subject to an examination by a digestion method with negative the meat has been found fit for human consumption following a post-mortem inspection carried out in accordance II.1.4 with Chapter II of Section I and Chapters VIII and IX of Section IV of Annex I to Regulation (EC) No 854/2004; II.1.5 (1) either [the carcass or parts of the carcass have been marked with a health mark in accordance with Chapter III of Section I of Annex I to Regulation (EC) No 854/2004;] (1) or [the packages of meat have been marked with an identification mark in accordance with Section I of Annex II to Regulation (EC) No 853/2004;] the meat satisfies the relevant criteria set out in Regulation (EC) No 2073/2005 on microbiological criteria for II.1.6 foodstuffs: II.1.7 the guarantees covering live animals and products thereof provided by the residue plans submitted in accordance with Directive 96/23/EC, and in particular Article 29 thereof, are fulfilled. the meat has been stored and transported in accordance with the relevant requirements of Section I of Annex III to II.1.8 Regulation (EC) No 853/2004 II.2. Animal Health attestation I, the undersigned official veterinarian, hereby certify, that the fresh meat described in Part I: II.2.1 has been obtained in the territory/ies with code:(2) which, at the date of issuing this certificate: [(a) has been free for 12 months from foot-and-mouth disease, rinderpest, African swine fever, (1) either classical swine fever, swine vesicular disease, and] (1) or $\hbox{\it [(a) (i)} \quad \hbox{has been free for 12 months from rinderpest, African swine fever, [foot-and-mouth disease] ('),}$ [classical swine fever] (1) and [swine vesicular disease] (1), and

[swine vesicular disease] (1), since

(EU) No, of ...

territory;

(ii) has been considered free from [foot-and-mouth disease] (1), [classical swine fever] (1) and

(b) during the last 12 months no vaccination against these diseases have been carried out and imports of domestic animals vaccinated against these diseases are not permitted in this

cases/outbreaks afterwards, and authorised to export this meat by Commission Regulation

.. (dd/mm/yyyy), and]

.....(dd/mm/yyyy), without having had

COUNTRY Model SUW

II. Health information			II.a. Certificate reference number	II.b.		
II.2.2		has been obtained from wild animals that were killed between				
			eeds 20 km from the borders of a country or pa his fresh meat into the Union,	rt thereof, which is not authorised during this		
	(b) in an area point II.2.1;		uring the last 60 days, there has been no	restrictions for the diseases referred to in		
II.2.3.A	has been obtained from animals which after killing were transported within 12 hours for chilling [to a collection centre, and immediately afterwards] (1) to an approved game-handling establishment around which, within a radiu of 10 km, there has been no case/outbreak of the diseases referred to in point II.2.1 during the previous 40 days of in the event of a case of disease, the preparation of meat for importation into the Union has been authorised on after removal of all meat, and the total cleaning and disinfection of the establishment under the control of an official veterinarian;					
(¹) (⁴) [II.2.3.B	has been obtained from carcasses on which the following test for classical swine fever was carried out and provided negative results:					
	(¹) either	(¹) either [virus isolation from blood (EDTA);]				
	(¹) or	[virus isc	lation from samples of	;]		
	(¹) or	[immuno	fluorescence for viral antigen on samples of	;]]		
II.2.4	has been obtained and prepared without contact with other meats not complying with the conditions required certificate.					
Notes						

This certificate is meant for fresh meat, excluding offal and minced meat, of wild animals belonging to the Suidae, Tayassuidae, or Tapiridae families that are killed or hunted in the wild.

 $Fresh\ meat\ means\ all\ animal\ parts\ fit\ for\ human\ consumption\ whether\ fresh,\ chilled\ or\ frozen.$

After importation, unskinned carcasses must be conveyed without delay to the processing establishment of destination.

Part I

- Box reference I.8: Provide the code of territory as appearing in Part 1 of Annex II to Regulation (EU) No 206/2010.
- Box reference I.11: Place of origin: name and address of the dispatch establishment.
- Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor must inform the BIP of entry into the Union.
- Box reference I.19: Use the appropriate HS code: 02.03, 02.08.90 or 05.04.
- Box reference I.20: Indicate total gross weight and total net weight.
- Box reference I.23: For containers or boxes, the container number and the seal number (if applicable) should be included.
- Box reference I.28: Nature of commodity: Indicate 'carcass-whole', 'carcass-side', 'carcass-quarters' or 'cuts'.
- Box reference I.28: Treatment type: If appropriate, indicate 'matured' or 'unskinned'. If frozen, indicate the date of freezing (mm/yy) of the cuts/pieces.
- Box reference I.28: Abattoir: any abattoir or game handling establishment.

	Model SU
II.a. Certificate reference number	II.b.
be authorised when obtained from animals killed third country, territory or part thereof referred to een adopted by the Union against imports of the provided when required in column 5 'SG' of Pare, in tests other than EDTA, the samples to be sample of at least one of the following lymph	2010. If or hunted either prior to the date of authorisation in boxes reference I.7 and I.8, or during a period his meat from this third country, territory or part to 1 of Annex II to Regulation (EU) No 206/2010, used are a sample of tonsil and of spleen plus nodes: retropharyngeal, parotid, mandibular or
Qualifica	ation and title:
Signatur	
J.g.m.	
	in Part 1 of Annex II to Regulation (EU) No 206// the authorised when obtained from animals killed third country, territory or part thereof referred to seen adopted by the Union against imports of the provided when required in column 5 'SG' of Part see, in tests other than EDTA, the samples to be to sample of at least one of the following lymph to be indicated. Qualification

	00	IIIITDV		Mode	I EQW				
_		UNTRY			I O Carrellia	ate reference		Veterinary certi	ricate to E
	1.1.	Consignor			i.z. Certific	ate reference	number	I.2.a.	
		Name Address			I.3. Central	Competent A	uthority		
_		Tel. No			I.4. Local C	ompetent Aut	hority		
me l	1.5	Consignee			I.6.				
li gu	1.5.	Name			1.0.				
čo Co		Address							
hed		Postal code							
patc		Tel. No							
Part I: Details of dispatched consignment	1.7.	Country ISO code	I.8. Region C of origin	ode	I.9. Country destina		SO ode	I.10. Region of destination	Code
)etai	1.11.	Place of origin			I.12.	l l			
-		Name	Approval number						
5		Address							
ł	1.13	. Place of loading		I.14. Date of	departure				
	1.15.	. Means of transport Aeroplane Sh	- I	I.16. Entry B	IP in EU				
			_						
		Road vehicle Oth							
		Identification: Documentary references:		1.17.					
	I.18	. Description of commodity				I.19. Comm	odity cod	de (HS code)	
							I.20. Q	uantity	
Ì	1.21	. Temperature of product					I.22. N	umber of packages	
		Ambient	Chiled		Frozen]			
İ	1.23	ldentification of container/s	eal number				I.24. T	ype of packaging	
	1.25	. Commodities certified for:							
		Human consumption							
	1.26			I.27. For import or admission into EU					
ŀ	1.28	. Identification of the commo	dities						
	10			val nu	mber establish	nments	_	Number	Net
	(\$	Scientific name) cor	mmodity Abattoir	Cı	utting plant	Cold store	0	f packages	weight

COUNTRY Model EQW

II. Health information II.a. Certificate reference number II.b. II.1. **Public Health Attestation** I, the undersigned official veterinarian, declare that I am aware of the relevant requirements of Regulations (EC) No 178/2002, (EC) No 852/2004, (EC) No 853/2004 and (EC) No 854/2004 and hereby certify that the meat of wild solipeds belonging to the subgenus Hippotigris (zebra) described in Part I was produced in accordance with those requirements, in particular Part II: Certification 11.1.1 the meat comes from (an) establishment(s) implementing a programme based on the HACCP principles in accordance with Regulation (EC) No 852/2004; II.1.2 the meat was obtained in compliance with Section IV of Annex III to Regulation (EC) No 853/2004; the meat fulfils the requirements of Regulation (EC) No 2075/2005 laying down specific rules on official controls for Trichinella in meat, in particular, has been subject to an examination by a digestion method with negative results; the meat has been found fit for human consumption following a post-mortem inspection carried out in accordance with Chapter II of Section I and Chapters VIII and IX of Section IV of Annex I to Regulation (EC) No 854/2004; II.1.4 [the carcass or parts of the carcass have been marked with a health mark in accordance with Chapter III of Section I of Annex I to Regulation (EC) No 854/2004;] Ithe packages of meat have been marked with an identification mark in accordance with Section I of (1) or Annex II to Regulation (EC) No 853/2004;] the meat satisfies the relevant criteria set out in Regulation (EC) No 2073/2005 on microbiological criteria for foodstuffs; the guarantees covering live animals and products thereof provided by the residue plans submitted in accordance with Directive 96/23/EC, and in particular Article 29 thereof, are fulfilled; 11.1.7 the meat has been stored and transported in accordance with the relevant requirements of Section I of Annex III to Regulation (EC) No 853/2004. II.2. **Animal Health attestation** I, the undersigned official veterinarian, hereby certify, that the fresh meat described in Part I: has been obtained from wild animals which after killing were transported within 12 hours for chilling [to a collection centre, and immediately afterwards] (1) to an approved game-handling establishment around which, within a radius of 10 km, there has been no case/outbreak of African horse sickness or glanders during the previous 40 days or, in the event of a case of such diseases, the preparation of meat for exportation to the Union has been authorised only after removal of all meat, and the total cleaning and disinfection of the establishment under the control of an official II.2.3 has been obtained and prepared without contact with other meats not complying with the requirements set out in this certificate Notes This certificate is meant for fresh meat, excluding offal and minced meat, of wild solipeds belonging to the subgenus Hippotigris Fresh meat means all animal parts fit for human consumption whether fresh, chilled or frozen. After importation, unskinned carcasses must be conveyed without delay to the processing establishment of destination.

COUNTRY		Model EQV
II. Health information	II.a. Certificate reference number	II.b.
provided. In case of unloading and relocement of the continuous provided. In case of unloading and relocement of the continuous provided in the continuous provided in the cuts/pieces. Box reference I.28: Nature of commodition of the cuts/pieces. Box reference I.28: Abattoir: any abatto Part II: (1) Keep as appropriate. (2) Dates. Imports of this meat shall not be a for importation into the Union of the this	te and address of the dispatch establishme or (railway wagons or container and lorries) ading, the consignor must inform the BIP of the Scode: 02.08.90 or 05.04. Weight and total net weight. Excess, the container number and the seal nuty: Indicate 'carcass-whole', 'carcass-side', opropriate, indicate 'matured' or 'unskinned in or game handling establishment. The stable of the seal of th	nt. In flight number (aircraft) or name (ship) is to be fentry into the Union. In the Union. In the Union
Official veterinarian		
Name (in capital letters):	Qualifica	ation and title:
Date:	Signatur	e:
Stamp:		

ANNEX III

Model TRANSIT/STORAGE

	COUNTRY	WoderTRA	NSII/STORAGE	Veterinary certificate to EU			
	I.1. Consignor		I.2. Certificate reference r	number I.2.a.			
	Name		I.3. Central Competent Au	uthority			
	Address						
ent	Tel. No		I.4. Local Competent Auth	nority			
ug	I.5. Consignee		I.6. Person responsible fo	or the consignment in EU			
onsi	Name		Name				
o p	Address		Address				
tche	Postal code		Postal code				
lispa	Tel. No		Tel. No				
Part I: Details of dispatched consignment	I.7. Country ISO of origin code	I.8. Region Code of origin	,	SO I.10. Region of Code destination			
Det	I.11. Place of origin		I.12. Place of destination				
art I:	Name	Approval number	Custom warehouse	Ship supplier			
ď	Address		Name Address	Approval number			
			Postal code				
	I.13. Place of loading		I.14. Date of departure				
	I.15. Means of transport		I.16. Entry BIP in EU	I.16. Entry BIP in EU			
	Aeroplane Ship	P Railway wagon					
	Road vehicle Othe	ır 🗌					
	Identification: Documentary references:		I.17. No. (s) of CITES	I.17. No. (s) of CITES			
	I.18. Description of commodity		I.19. Commo	odity code (HS code)			
				I.20. Quantity			
	I.21. Temperature of product			I.22. Number of packages			
	Ambient	Chiled	Frozen	, , , , , , , , , , , , , , , , , , ,			
	I.23. Identification of container/se	al number		I.24. Type of packaging			
	I.25. Commodities certified for: Human consumption						
	I.26. For transit through EU to 3 ro	d Country	I.27.				
	3rd country	ISO code					
	I.28. Identification of the commod	lities					
	Species Nature of (Scientific name) commodit		number establishments	Number Net of packages weight			
	(Salahana nama)	Abattoir	Cutting manufacturing plant/ plant	o. pasiagos moigrit			

COUNTRY	Model TRANSIT/STORAGE

	II.	Health information	II.a. Certif	icate reference number	II.b.		
	II.1.	Animal Health Attestati	on				
		I, the undersigned official	veterinarian, hereby	y certify, that the fresh meat descri	bed in Part I:		
_				egion authorized for imports into the Union as laid down in Part 1 of Annex II to Regulation ime of slaughter, and			
Part II: Certification			[OVI] [POR] [EQU]		the animal health attestation in the model [(1) in Part 2 of Annex II to Regulation (EU)		
Part II: C				e slaughtered and processed on mm/yyyy) and	(dd/mm/yyyy) (²).		
	Notes				0 (D) 11 07/70/70 (
		ificate is meant for transit a i meat, including minced m	-	dance with Article 12(4) or Article 1	3 of Directive 9///8/EC of:		
	(1)	_		and <i>Bison</i> species and their cross	-breeds) (Model 'BOV'):		
	(2)		,	stic caprine animals (Capra hircus			
	(3)	domestic porcine animals	•	. , , .	, (
		meat, excluding minced m		,			
	(4)			sinus and their cross-breeds) (Mod	del 'EQU'):		
	` '	meat, excluding offal and		, (,,		
	(5)	farmed non-domestic ani	mals of the order Art		ls (including <i>Bison</i> and <i>Bubalus</i> species and e families Rhinocerotidae and Elephantidae.		
	(6)				(including <i>Bison</i> and <i>Bubalus</i> species and le families Rhinocerotidae and Elephantidae		
	(7)	farmed non-domestic ani	mals belonging to th	ne Suidae, Tayassuidae, or Tapirida	ae families (Model 'SUF');		
	(8)	wild non-domestic anima	Is belonging to the S	Suidae, Tayassuidae, or Tapiridae f	amilies (Model 'SUW');		

(9) wild solipeds belonging to the subgenus *Hippotigris* (zebra) (Model 'EQW').

Fresh meat means all animal parts fit for human consumption whether fresh, chilled or frozen.

COUNTRY	Model TRANSIT/STORAGE
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II.	Health information	II.a. Certificate reference number	II.b.					
Pa	Part I:							
	Box reference I.8: Provide the code of te Box reference I.11: Place of origin: name Box reference I.12: Address (and approvor ship chandler shall be included. Box reference I.15: Registration numbe provided. In case of unloading and reload Box reference I.19: Use the appropriate Box reference I.20: Indicate total gross of Box reference I.23: For containers or both Box reference I.28: Nature of commodity Box reference I.28: Treatment type: If from the III: Keep as appropriate. Date or dates of slaughter. Imports of the date of authorisation for exportation to the	erritory as appearing in Part 1 of Annex II to Rele and address of the dispatch establishment. val number if known) of the warehouse in a free or (railway wagons or container and lorries), fligating, the consignor must inform the BIP of en HS code: 02.01, 02.02, 02.03, 02.04, 02.05, 02.04, and total net weight. oxes, the container number and the seal number. Indicate 'carcass-whole', 'carcass-side', 'carboxen, indicate the date of freezing (mm/yy) of the seal of the container oxes. The container oxes are union of the third country, territory or part the oxes been adopted by the Union against imports.	ezone, free warehouse, customs warehouse ght number (aircraft) or name (ship) is to be try into the Union. 12.06, 02.08.90, 02.09, 05.04 or 15.02. er (if applicable) should be included. cass-quarters', 'cuts', or 'minced meat'. he cuts/pieces.					
Off	Official veterinarian							
	Name (in capital letters):	Qualification	and title:					
	Date:	Signature:						
	Stamp:							

ANNEX IV

ANIMALS REFERRED TO IN ARTICLE 1(1)(b)

PART 1

Lists of third countries, territories or parts thereof

SECTION 1

Parts of third countries or territories referred to in Article 7(2)

▼<u>M1</u>

Country/territory	Code of part of the country/territory	Description of part of the country/territory		
US – United States	US-A	The State of Hawaii (1)		
(1) Suspended from 5 May 2010.				

▼<u>C1</u>

PART 2

Tables of animals and the corresponding model veterinary certificates

Table 1				
'QUE': Model of veterinary certificate for consignments of queen bees and queen bumble bees (Apis mellifera and Bombus spp.),				
'BEE': Model of veterinary certificate for consignments of colonies of bumble bees (<i>Bombus</i> spp.				
Order Family Genera/species				
Hymenoptera	Apidae Apis mellifera, Bombus			

			el QUE			
		UNTRY	Veterinary certificate to El			
	1.1.	Consignor	I.2. Certificate reference number I.2.a.			
		Name	I.3. Central Competent Authority			
		Address	I.4. Local Competent Authority			
		Tel. No	1.4. Local Competent Authority			
ent	1.5.	Consignee	1.6.			
muk		Name				
nsig		Address				
o p		Postal code				
che		Tel. No				
Part I: Details of dispatched consignment	1.7.	Country ISO I.8. Region Code of origin code	I.9. Country of ISO I.10. Region of Code destination code destination			
ilso	1.11	Place of origin	I.12.			
l: Deta		Name Approval number Address				
Part		Name Approval number Address				
		Name Approval number Address				
	1.13	. Place of loading	I.14. Date of departure time of departure			
		Address Approval number				
	I.15	Means of transport Aeroplane Ship Railway wagon	I.16. Entry BIP in EU I.17. No(s) of CITES			
		Road vehicle Other Identification:				
	I.18	Documentary references: Description of commodity	I.19. Commodity code (HS code) 01.06.90			
			I.20. Quantity			
	I.21		I.22. Number of packages			
	1.23	. Identification of container/seal number	1.24.			
	I.25. Commodities certified for: Breeding					
	1.26		I.27. For import or admission into EU			
	1.28. Identification of the commodities					
			ication Identification tem number			

	COUNT	RY			Model QUE	
	II.	Health i	information	II.a. Certificate reference number	II.b.	
	II.1. Animal Health attestation:					
		I, the un	ndersigned, hereby certify	r, that the animals referred to in Part I o	f this certificate meet the following requirements:	
Ę		II.1.1 they come from the territory with code:(!) in which, American foulbrood, the small hive beetle (Aet tumida) and the Tropilaelaps mite (Tropilaelaps spp.) are notifiable diseases/pests.				
catio		II.1.2	I.1.2 they:			
ertific			(a) come from a breeding	ng apiary, which is supervised and con	trolled by the competent authority;	
Part II: Certification		(b) come from an area which is not subject to any restrictions associated with an occurrence of American foulbrood and where no such occurrence has taken place within at least 30 days prior to the issuance of the presen certificate. Where an outbreak of American foulbrood has occurred previously, all hives within a radius of three kilometres have been checked by the competent authority and all infected hives burned or treated and inspected to the satisfaction of the said competent authority within 30 days following the last recorded case:				
	(c) are from hives or come from hives or colonies (in the case of bumble bees) from which samples of the have been tested in the last 30 days for American foulbrood as laid down in the OIE Manual of Diagnotand Vaccines for terrestrial Animals with negative results;					
					ject to any restrictions associated with the occurrence up, and where these infestations are absent;	
					of bumble bees), which were inspected immediately disease including infestations affecting bees;	
					bees and packaging do not contain the small hive infestations, in particular <i>Tropilaelaps</i> spp., affecting	
			diseased bees or brood-	combs, and all precautions have been	and food are new and have not been in contact with taken to prevent contamination with agents causing	
	Notes					
	Part I:					
	 Box reference I.20: Number of queen bees (Apis mellifera and Bombus spp.). Each queen bee may be accompanied by a ma of 20 attendants. 				Each queen bee may be accompanied by a maximum	
	Part II:					
	(¹) Code of the territory as it appears in Part 1 of Annex II or Section 1 of Part 1 of Annex IV to Regulation (EU) No 206/2010. Official veterinarian /Official inspector				f Annex IV to Regulation (EU) No 206/2010.	
	Name (in capital letters):		Qua	alification and title:		
		Date:		Sign	nature:	
		Stamp:				

	Model BEE COUNTRY Veterinary certificate to EU							
		Consignor	Veterinary certificate to EU 1.2. Certificate reference number 1.2.a.					
	1.1.	Name	i.z. Certifica	ale relerence i	number	1.2.a.		
		Address	I.3. Central	Competent A	uthority			
		Tel. No	I.4. Local C	ompetent Aut	hority			
	1.5	Consignee	I.6.					
nent	1.5.			1.0.				
Part I: Details of dispatched consignment		Name Address						
ed c		Postal code						
atch	17	Tel. No	Dagion Code	10 Country	, of 10	20	I 10. Degion of	Codo
of dispa	1.7.		Region Code of origin	I.9. Country destina		SO I	I.10. Region of destination	Code
ails	1.11	. Place of origin		I.12.				
l: Deta		Name Appr Address	roval number					
Part		Name Appr Address	roval number					
		Name Appr Address						
	I.13. Place of loading Address Approval number			I.14. Date of departure time of departure				
	I.15. Means of transport Aeroplane			I.16. Entry B	IP in EU			
				I.17. No(s) of CITES				
	I.18	. Description of commodity			I.19. Commo	odity cod	de (HS code) 0	1.06.90
						I.20. Q	uantity	
	1.21	ı.				I.22. N	umber of packages	3
	I.23. Identification of container/seal number			1.24.				
	I.25. Commodities certified for:							
	Breeding							
	1.26.			I.27. For imp	ort or admissi	on into E	:U [
	I.28. Identification of the commodities							
				ication tem			Identification number	

	COUNT	RY		Model BEE	
	II.	Health information	II.a. Certificate reference number	II.b.	
	II.1. Animal Health attestation:				
		I, the undersigned, hereby certify	y that:		
		II.1.1			
ication			ombus spp.) referred to in Part I of this certificate a recognised establishment which is supervise		
Part II: Certification	 (b) the establishment referred to in Part I of this certificate was inspected immediately prior to dispatch an bumble bees and breeding stock show no clinical signs or suspicion of disease including infestations affer bees; 				
Pa	(c) all colonies for import into the Union have undergone detailed examination to ensure that all bumble bee broodstock and packaging do not contain the small hive beetle (Aethina tumida) or its eggs and larvae or other infestations in particular Tropilaelaps spp., affecting bees;				
			ontainers, accompanying products and food a -combs, and all precautions have been taken to of bees.		
	Notes				
	Part I:				
	Box reference I.20: Number of containers of bumble bees (<i>Bombus</i> spp.), each containing a colony of a maximum of 200 bumble bees. Official veterinarian /Official inspector				
		Name (in capital letters):	Qualification	n and title:	
		Date:	Signature:		
		Stamp:			

ANNEX V

Explanatory notes for completing the veterinary certificates

(referred to in Article 18)

(a) Veterinary certificates shall be issued by the exporting third country, based on the models set out in Part 2 of Annexes I, II and IV and Annex III according to the layout of the model that corresponds to the live animals/fresh meat concerned.

They shall contain, in the numbered order that appears in the model, the attestations that are required for any third country and, as the case may be, those supplementary guarantees that are required for the exporting third country or part thereof.

If the Member State of destination imposes, for the live animals/fresh meat concerned, additional certification requirements, attestations to certify that those requirements are fulfilled shall also be incorporated in the original form of the veterinary certificate.

- (b) Where the model certificate states that certain statements shall be kept as appropriate, statements which are not relevant, may be crossed out and initialled and stamped by the certifying officer, or completely deleted from the certificate
- (c) A separate and unique certificate must be provided for the live animals/fresh meat that are exported from a territory or territories of the same exporting country appearing in columns 2 and 3 of Part 1 of Annex I, II or IV which are consigned to the same destination and transported in the same railway wagon, lorry, aircraft or ship.
- (d) The original of each certificate shall consist of a single sheet of paper, or, where more text is required it must be in such a form that all sheets of paper required are part of an integrated whole and indivisible.
- (e) The veterinary certificate shall be drawn up in at least one of the official languages of the Member State of the border inspection post of introduction of the consignment into the Union and of the Member State of destination. However, those Member States may authorise the certificate to be drawn up in the official language of another Member State, and accompanied, if necessary, by an official translation.
- (f) If for reasons of identification of the items of the consignment (schedule in point 1.28 of the model veterinary certificate), additional sheets of paper are attached to the certificate, those sheets of paper shall also be considered as forming part of the original of the certificate by the application of the signature and stamp of the certifying officer, on each of the pages.
- (g) When the certificate, including additional schedules referred to in (f), comprises more than one page, each page shall be numbered, (page number) of (total number of pages), at the end of the page and shall bear the certificate reference number that has been designated by the competent authority at the top of the pages.
- (h) The original of the certificate must be completed and signed by an official veterinarian or by another designated official inspector where this is provided for in the model veterinary certificate. In the case of live animals, the certificate must be completed and signed within 24 hours prior to loading of the consignment for introduction into the Union. The competent authorities of the exporting third country shall ensure that rules of certification equivalent to those laid down in Directive 96/93/EC (¹) are followed.

The colour of the signature shall be different from that of the printing. This requirement also applies to stamps other than those embossed or watermarked.

(i) The certificate reference number referred to in boxes I.2 and II.a. must be issued by the competent authority.