CHAPTER 38

MODEL ANIMAL HEALTH CERTIFICATE FOR ENTRY INTO THE UNION OF DOGS, CATS AND FERRETS (MODEL ‘CANIS-FELIS-FERRETS’)

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| **COUNTRY URUGUAY** | | | | **Animal health certificate to the EU** | | | |
| **Part I: Description of consignment** | **I.1 Consignor/Exporter**  Name Address  Country **URUGUAY** ISO country code **UY** | | | **I.2 Certificate reference** | | **I.2a IMSOC reference** | |
| **QR CODE** | |
| **I.3 Central Competent Authority** | |
| **I.4 Local Competent Authority** | |
| **I.5 Consignee/Importer**  Name Address  Country **URUGUAY** ISO country code **UY** | | | **I.6 Operator responsible for the consignment**  Name Address  Country ISO country code | | | |
| **I.7 Country of origin URUGUAY** ISO country code **UY** | | | **I.9 Country of destination** ISO country code | | | |
| **I.8 Region of origin URUGUAY** Code **UY** | | | **I.10 Region of destination** Code | | | |
| **I.11 Place of dispatch**  Name Registration/Approval No Address  Country **URUGUAY** ISO country code **UY** | | | **I.12 Place of destination**  Name Registration/Approval No  Address  Country ISO country code | | | |
| **I.13 Place of loading** | | | **I.14 Date and time of departure** | | | |
|  | **I.15 Means of transport**   * Aircraft **□** Vessel * Railway **□** Road vehicle Identification | | | **I.16 Entry Border Control Post** | | | |
| **I.17 Accompanying documents**  Type Code  Country ISO country code  Commercial document reference | | | |
| **I.18** | **Transport conditions** | **□** Ambient | | **□** Chilled | | **□** Frozen |
| **I.19 Container number/Seal number**  Container No Seal No | | | | | | |
| **I.20** | **Certified as or for** | | | | | |
| * Further keeping   + Confined establishment   + Quarantine establishment   **□** Other | | | | | | |
| **I.21 □ For transit**  Third country ISO country code | | | **I.22 □ For internal market** | | | |
| **I.23** | | | |

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| **I.24 Total number of packages** | | | **I.25 Total quantity** | | | | **I.26 Total net weight/gross weight (kg)** | | |
| **I.27 Description of consignment** | | | | | | | | | |
| CN code | Species | Subspecies/Category | | Sex | Identification system  Nature of commodity | Identification number | | Age  Test | Quantity |

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| **Part II: Certification** | **II. Health information** | **II.a Certificate reference** | **II.b IMSOC reference** |
| I, the undersigned official veterinarian hereby certify that the animals described in Part I:   * 1. come from a country, territory or zone thereof with code: - (1) which, on the date of issue of this certificate is authorised for the entry into the Union of dogs, cats and ferrets and is listed in Part 1 of Annex VIII to Commission Implementing Regulation (EU) 2021/404;   *(2)(3)either* [II.2. have been dispatched directly from the establishment of origin to the Union without passing through any other establishment];  *(2)(3)or* [II.2. have undergone one single assembly operation in the country, territory or zone thereof of origin which took place for not more than 6 days in an establishment fulfilling the following requirements:   * + - it is approved for conducting assembly operations of dogs, cats and ferrets by the competent authority in the third country or territory in accordance with Article 10 of Commission Delegated Regulation (EU) 2019/2035;     - it has a unique approval number assigned by the competent authority of the third country or territory;     - it is listed for that purpose by the competent authority of the third country or territory of dispatch, including the information set out in Article 21 of Delegated Regulation (EU) 2019/2035;     - it complies with the record keeping requirements provided for in point (a)(iv) of Article 73(2) of Delegated Regulation (EU) 2020/692.]   (3)[II.3. have been loaded for dispatch to the Union on / / (dd/mm/yyyy)(4) in a means of transport which was cleaned and disinfected prior to loading with a disinfectant authorised by the competent authority in the third country or territory and constructed in such a way that:   * + - animals cannot escape or fall out;     - visual inspection of the space where animals are kept is possible;     - the escape of animal excrements, litter or feed is prevented or minimized.]   II.4 have been subjected with negative result to a clinical inspection, carried out by an official veterinarian in the third country, territory or zone thereof of origin within 48 hour period prior to loading for dispatch to the Union for the detection of signs indicative of the occurrence of diseases, including the relevant listed diseases referred to in Annex 1 of Delegated Regulation (EU) 2020/692 and emerging diseases.  *(2)either* [II.5. are destined for direct entry into the Member State of destination to be isolated in:  *(2)either* [a confined establishment;]]  *(2)or* [an approved quarantine establishment;]] | | |

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*(2)or* [II.5. were at least 12 weeks old at the time of vaccination against rabies and at least 21 days have elapsed since the completion of the primary anti-rabies vaccination(5) carried out in accordance with the validity requirements set out in Annex III to Regulation (EU) No 576/2013 of the European Parliament and of the Council, and any subsequent revaccination was carried out within the period of validity of the preceding vaccination(6), and

*(2)either* [they come from, and in case of transit are scheduled to transit through, a territory or third country listed in Annex II to Commission Implementing Regulation (EU) No 577/2013 and details of the current anti-rabies vaccination are provided in columns 1 to 7 in the table below;]]

*(2)or* [they come from or are scheduled to transit through a territory or third country not listed in Annex II to Commission Implementing Regulation (EU) No 577/2013, and

details of the current anti-rabies vaccination are provided in columns 1 to 7 in the table below, and

- a rabies antibody titration test(7), carried out on a blood sample taken by the veterinarian authorised by the competent authority not less than 30 days after the preceding vaccination and at least three months prior to the date of issue of this certificate, proved an antibody titre equal to or greater than 0,5 IU/ml(8) and any subsequent revaccination was carried out within the period of validity of the preceding vaccination, and the date of sampling for testing the immune response are provided in column 8 in the table below:]]

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| **Transponder** | | **Date of vaccination [dd/mm/yyyy]** | **Name and manufacturer of vaccine** | **Batch number** | **Validity of vaccination** | | **Date of blood sampling [dd/mm/yyyy]** |
| **Alphanumeric code of the animal** | **Date of implantation and/or reading (9) [dd/mm/yyyy**  **]** |
| **From dd/mm/yyyy**  **]** | **To dd/mm/yyyy**  **]** |
| 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
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*(2)either* [II.6. the consignment includes dogs destined for a Member State listed in the Annex to Commission Implementing Regulation (EU) 2018/878 and those dogs have been treated against infestation with *Echinococcus multilocularis*, and the details of the treatment carried out by the administering veterinarian in accordance with point 2 of Annex XXI to Delegated Regulation (EU) 2020/692(10) (11) are provided in the table below

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| **Transponder or tattoo. Alphanumeric code of the dog** | **Anti-Echinococcus treatment** | | **Administering veterinarian** |
| **Name and manufacturer of the product** | **Date [dd/mm/yyyy] and time of treatment [00:00]** | **Name in capitals, stamp and signature** |
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*(2)or* [II.6. the dogs have not been treated against infestation with *Echinococcus multilocularis.*]

*(2)or* [II.6. the dogs are destined for direct entry into the Member State of destination to be isolated in :

(1)either [a confined establishment.]]

(1)or [an approved quarantine establishment.]]

**Notes:**

This certificate is intended for commercial entries into the Union of dogs, cats and ferrets, including when they are destined to a confined establishment or to an approved quarantine establishment and when the Union is not the final destination of the animals and for entry into the Union of dogs, cats and ferrets moved in accordance with Article 5(4) of Regulation (EU) No 576/2013 of the European Parliament and of the Council.

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

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

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|  | **Part I:**  Box I.20: Certified as or for: indicate   * "Further keeping" where dogs, cats or ferrets are moved in accordance with Title V of Part II of Delegated Regulation (EU) 2020/692; * Confined establishment: as defined in Article 4(48) of Regulation (EU) 2016/429 of the European Parliament and of the Council; * Approved quarantine establishment: as defined in Article 3(9) of Commission Delegated Regulation (EU) 2020/688 * "others" where dogs (*Canis lupus familiaris*), cats (*Felis silvestris catus*) or ferrets (*Mustela putorius furo*) are moved in accordance with Article 5(4) of Regulation (EU) No 576/2013 of the European Parliament and of the Council.   **Part II:**  (1) Code of the zone as it appears in Column 2 of Part 1 of Annex VIII to Implementing Regulation (EU) 2021/404.  (2) Keep as appropriate.  (3) Not applicable to the movement of dogs, cats and ferrets other than non-commercial movements kept as pet animals in households that cannot be carried out in accordance with the conditions laid down in Article 245(2) or Article 246(1) and (2) of Regulation (EU) 2016/429.  (4) Date of loading: it cannot be a date prior to the date of authorisation of the zone for entry into the Union, or a date in a period when restriction measures have been adopted by the Union against entries of these animals from the zone.  (5) Any revaccination must be considered a primary vaccination if it was not carried out within the period of validity of a previous vaccination.  (6) A certified copy of the identification and vaccination details of the animals concerned shall be attached to the certificate.  (7) The rabies antibody titration test referred to in point II.5:   * must be carried out on a sample collected by a veterinarian authorised by the competent authority, at least 30 days after the date of vaccination and three months before the date of import*;* * must measure a level of neutralising antibody to rabies virus in serum equal to or greater than 0,5 IU/ml; * must be performed by an official laboratory; * does not have to be renewed on an animal, which following that test with satisfactory results, has been revaccinated against rabies within the period of validity of a previous vaccination.   A certified copy of the official report from the official laboratory on the result of the rabies antibody test referred to in point II.5. shall be attached to the certificate.  (8) By certifying this result, the official veterinarian confirms that he has verified, to the best of his ability and where necessary with contacts with the laboratory indicated in the report, the authenticity of the laboratory report on the results of the antibody titration test referred to in point II.5.  (9) In conjunction with footnote (6), the marking of the animals concerned by the implantation of a transponder must be verified before any entry is made in this certificate and must always precede any vaccination, or where applicable, testing carried out on those animals. |
|  | (10) The treatment against infestation with *Echinococcus multilocularis* referred to in point II.6 must:   * be administered by a veterinarian within a period of not more than 48 hours and ending not less than 24 hours before the time of the scheduled entry of the dogs into one of the Member States or parts thereof listed in the Annex to Commission Implementing Regulation (EU) 2018/878; * consist of an approved medicinal product which contains the appropriate dose of praziquantel or pharmacologically active substances, which alone or in combination, have been proven to reduce the burden of mature and immature intestinal forms of *Echinococcus multilocularis* in the host species concerned.   (11) The table referred to in point II.6 must be used to document the details of a further treatment if administered after the date the certificate was signed and prior to the scheduled entry into one of the Member States or parts thereof listed in the Annex to Commission Implementing Regulation (EU) 2018/878. |
|  | **Official veterinarian** |

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|  | Name (in capital letters)  Date Qualification and title  Stamp Signature |