Model animal health certificate for the non-commercial movement into a Member State from a territory or third country of dogs, cats or ferrets in accordance with Article 5(1) and (2) of Regulation (EU) No 576/2013

CO	OUNTRY:	Veterinary certificate to EU				
P	I.1. Consignor	I.2. Certificate reference No I.2.a.				
a	Name Address	I.3. Central competent authority				
r t	Tel.	I.4. Local competent authority I.6. Person responsible for the consignment in the EU				
I : D e t a	I.5. Consignee Name Address Postal code Tel.					
i l s	I.7. Country of origin ISO code I.8. Region of origin Code	I.9. Country of ISO code I.10 Region of Code destination				
o f d i s	I.11. Place of origin	I.12. Place of destination				
a t c h e d c o n s i g n	I.13. Place of loading	I.14. Date of departure				
e n t						
	I.15. Means of transport	I.16. Entry BIP in EU				
		I.17. No.(s) of CITES				
	I.18. Description of commodity	I.19. Commodity code (HS code) 010619				
		I.20. Quantity				
	I.21. Temperature of products	I.22. Total number of packages				
	I.23. Seal/Container No	I.24. Type of packaging				
	I.25. Commodities certified for: Pets •					

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I.26. For transit to	o 3 rd Cou	ıntry			I.27. Fo	r import or admission into	EU
I.28. Identification of the commodities							
Species (Scientific name)	Sex	Colour	Breed	Identification no	umber	Identification system	Date of birth [dd/mm/yyyy]

COUNTRY

Non-commercial movement into a Member State from a territory or third country of dogs, cats or ferrets in accordance with Article 5(1) and (2) of Regulation (EU) No 576/2013

	II. Health i	nformation	II.a. Certificate reference No	o II.b.				
Dowt			narian ⁽¹⁾ /veterinarian authorised b(insert name of territory	or third country) certify that:				
Part II: Ce rtif icat	<u>Purpose/</u> II.1.	the owner to carry ou supported by evidence ⁽ the natural person who						
ion	(1) 224, 22	non-commercial mover	movement that aims at their sale or a transfer of ownership, and during the ent will remain under the responsibility of					
	⁽¹⁾ either ⁽¹⁾ or		has authorisation in writing from the owner to carry out the non-commercial s on behalf of the owner;]					
	(l)or	movement of the anima	gnated by a carrier contracted by the owner to carry out the non-commercial is on behalf of the owner;]					
	(1) either [II.2. (1) or [II.2.	the animals described in Box I.28 are moved in a number of five or less;] the animals described in Box I.28 are moved in a number of more than five, are more than six months old and are going to participate in competitions, exhibitions or sporting events or in training for those events, and the owner or the natural person referred to in point II.1 has provided evidence ⁽³⁾ that the animals are registered						
	⁽¹⁾ either ⁽¹⁾ or	[to attend such event;] [with an association organising such events;]						
	1		- 13					
	(1) either [II.3.	Attestation of rabies vaccination and rabies antibody titration test: (1) either [II.3. the animals described in Box I.28 are less than 12 weeks old and have not received an antivaccination, or are between 12 and 16 weeks old and have received an anti-rabies vaccination, days at least have not elapsed since the completion of the primary vaccination against rabies out in accordance with the validity requirements set out in Annex III to Regulation (Elapsed S76/2013 ⁽⁴⁾ , and						
		Annex II to destination in	r third country of provenance of the ar Implementing Regulation (EU) No dicated in Box I.5 has informed the pub- into its territory, and they are accompan	577/2013 and the Member State of blic that it authorises the movement of				
	⁽¹⁾ either	that from bir	leclaration ⁽⁵⁾ of the owner or the natural th until the time of the non-commercia wild animals of species susceptible to ral	il movement the animals have had no				
	⁽¹⁾ or	before their b	on whom they still depend, and it can b irth an anti-rabies vaccination which co ex III to Regulation (EU) No 576/2013;	omplied with the validity requirements				
	(1) or/and [II.3.	and at least 21 days carried out in accordan	n Box I.28 were at least 12 weeks old at the time of vaccination against rabies have elapsed since the completion of the primary anti-rabies vaccination (a) are with the validity requirements set out in Annex III to Regulation (EU) No esequent revaccination was carried out within the period of validity of the original properties of the original properties and the period of validity of the original properties are the period of validity or the original properties are the period of validity or the original properties are the period of validity or the original properties are the period of validity or the original properties are the period of validity or the original properties are the period					
	⁽¹⁾ either	II to Impleme third country a territory o Regulation (I	escribed in Box I.28 come from a terrisonting Regulation (EU) No 577/2013, edisted in Annex II to Implementing Regulation at third country other than those I (EU) No 577/2013 in accordance with potential potential (2013), and the details of the current a w;]	either directly, through a territory or a gulation (EU) No 577/2013 or through isted in Annex II to Implementing oint (c) of Article 12(1) of Regulation				
	⁽¹⁾ or	territory or the (EU) No 577	described in Box I.28 come from, or hird country other than those listed in A /2013 and a rabies antibody titration the veterinarian authorised by the competent	Annex II to Implementing Regulation test ⁽⁸⁾ , carried out on a blood sample				

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table below 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⁽⁹⁾ and any subsequent revaccination was carried out within the period of validity of the preceding vaccination⁽⁶⁾, and the details of the current anti-rabies vaccination and the date of sampling for testing the immune response are provided in the table below:

Transponder or tattoo					Validity of vaccination		
Alphanumeri c code of the animal	Date of implantation and/or reading ⁽¹⁰⁾ [dd/mm/yyyy]	Date of vaccination [dd/mm/yyyy]	Name and manufacturer of vaccine	Batch number	From [dd/mm/ yyyy]	to [dd/mm/ yyyy]	Date of the blood sampling [dd/mm/yyyy]

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Attestation of anti-parasite treatment:

(1) either [II.4.

the dogs described in Box I.28 are destined for a Member State listed in Annex I to Commission Delegated Regulation (EU) No 1152/2011 and have been treated against *Echinococcus multilocularis*, and the details of the treatment carried out by the administering veterinarian in accordance with Article 7 of Commission Delegated Regulation (EU) No 1152/2011⁽¹¹⁾⁽¹²⁾⁽¹³⁾ are provided in the table below.]

(1) or [II.4. the dogs described in Box I.28 have not been treated against Echinococcus multilocularis(11).]

Transponder or		chinococcus eatment	Administering veterinarian
tattoo number of the dog	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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Notes

- (a) This certificate is meant for dogs (*Canis lupus familiaris*), cats (*Felis silvestris catus*) and ferrets (*Mustela putorius furo*).
- (b) This certificate is valid for 10 days from the date of issue by the official veterinarian until the date of the documentary and identity checks at the designated Union travellers' point of entry (available at http://ec.europa.eu/food/animal/liveanimals/pets/pointsentry_en.htm).

In the case of transport by sea, that period of 10 days is extended by an additional period corresponding to the duration of the journey by sea.

For the purpose of further movement into other Member States, this certificate is valid from the date of the documentary and identity checks for a total of four months or until the date of expiry of the validity of the anti-rabies vaccination or until the conditions relating to animals less than 16 weeks old referred to in point II.3 cease to apply, whichever date is earlier. Please note that certain Member States have informed that the movement into their territory of animals less than 16 weeks old referred to in point II.3 is not authorised. You may wish to inquire at http://ec.europa.eu/food/animal/liveanimals/pets/index_en.htm.

Part I:

Box I.5: Consignee: indicate Member State of first destination.

Box I.28: Identification system: select of the following: transponder or tattoo.

Identification number: indicate the transponder or tattoo alphanumeric code.

Date of birth/breed: as stated by the owner.

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Part II:

- (1) Keep as appropriate.
- The declaration referred to in point II.1 shall be attached to the certificate and comply with the model and additional requirements set out in Part 3 of Annex IV to Implementing Regulation (EU) No 577/2013.
- The evidence referred to in point II.1 (e.g. boarding pass, flight ticket) and in point II. 2 (e.g. receipt of entry to the event, proof of membership) shall be surrendered on request by the competent authorities responsible for the checks referred to in point (b) of the Notes.
- Any revaccination must be considered a primary vaccination if it was not carried out within the period of validity of a previous vaccination.
- The declaration referred to in point II.3.2 to be attached to the certificate complies with the format, layout and language requirements laid down in Parts 1 and 3 of Annex I to Implementing Regulation (EU) No 577/2013
- (6) A certified copy of the identification and vaccination details of the animals concerned shall be attached to the certificate.
- The third option is subject to the condition that the owner or the natural person referred to in point II.1 provides, on request by the competent authorities responsible for the checks referred to in point (b), a declaration stating that the animals have had no contact with animals of species susceptible of rabies and remain secure within the means of transport or the perimeter of an international airport during the transit through a territory or a third country other than those listed in Annex II to Implementing Regulation (EU) No 577/2013. This declaration shall comply with the format, layout and language requirements set out in Parts 2 and 3 of Annex I to Implementing Regulation (EU) No 577/2013.
- (8) The rabies antibody titration test referred to in point II.3.1:
 - 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 a laboratory approved in accordance with Article 3 of Council Decision 2000/258/EC (list of approved laboratories available at http://ec.europa.eu/food/animal/liveanimals/pets/approval_en.htm);
 - 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 approved laboratory on the results of the rabies antibody test referred to in point II.3.1 shall be attached to the certificate.

- (9) 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.3.1.
- In conjunction with footnote (6), the marking of the animals concerned by the implantation of a transponder or by a clearly readable tattoo applied before 3 July 2011 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.
- The treatment against *Echinococcus multilocularis* referred to in point II.4 must:
 - be administered by a veterinarian within a period of not more than 120 hours and 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 Annex I to Delegated Regulation (EU) No 1152/2011;
 - 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.
- The table referred to in point II.4 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 Annex I to Delegated Regulation (EU) No 1152/2011.
- The table referred to in point II.4 must be used to document the details of treatments if administered after the date the certificate was signed for the purpose of further movement into other Member States described in point (b) of the Notes and in conjunction with footnote (11).

Official veterinarian/Authorised veterinarian						
Name (in capital letters):	Qualification and title:					
Address						
Telephone:						
Date:	Signature:					
Stamp:						
Endorsement by the competent authority (not necessary when	the certificate is signed by an official veterinarian)					
Name (in capital letters):	Qualification and title:					
Address						
Telephone:						
Date:	Signature:					
Stamp:						
Official at the travellers' point of entry (for the purpose of further movement into other Member States)						
Name (in capital letters):	Title:					
Address						
Telephone:						
E-mail address:						
Date of completion of the documentary and identity che	cks: Signature: Stamp:					

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