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COMMISSION IMPLEMENTING DECISION

of 21 October 2013

laying down the list of territories and third countries authorised for imports of dogs, cats and ferrets and the model health certificate for such imports

(notified under document C(2013) 6721)

(Text with EEA relevance)

(2013/519/EU)

(OJ L 281, 23.10.2013, p. 20)

Amended by:

		(Official Journal			
		No	page	date		
► <u>M1</u>	Commission Implementing Decision (EU) 2017/98 of 18 January 2017	L 16	37	20.1.2017		

►<u>B</u>

COMMISSION IMPLEMENTING DECISION

of 21 October 2013

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(notified under document C(2013) 6721)

(Text with EEA relevance)

(2013/519/EU)

Article 1

List of territories or third countries from which dogs, cats or ferrets are authorised to be imported in accordance with Directive 92/65/EEC

1. Consignments of dogs, cats or ferrets which are subject to the provisions of Directive 92/65/EEC shall only be imported into the Union provided that the territories or third countries they come from and any territories or third countries they transit are included in one of the lists set out in:

(a) Annex I to Decision 2004/211/EC;

(b) Part 1 of Annex II to Regulation (EU) No 206/2010;

(c) Annex II to Implementing Regulation (EU) No 577/2013.

2. By way of derogation from paragraph 1, consignments of dogs, cats or ferrets destined for bodies, institutes and centres approved in accordance with Directive 92/65/EEC shall only be imported into the Union provided that the territories or third countries they come from and any territories or third countries they transit are included in the list referred to in paragraph 1(c).

Article 2

Animal health certificate for imports from territories or third countries

Member States shall only authorise imports of dogs, cats or ferrets, which comply with the following conditions:

- (a) they are accompanied by an animal health certificate drawn up in accordance with the model set out in Part 1 of the Annex and completed and signed by an official veterinarian in accordance with the explanatory notes set out in Part 2 of the Annex;
- (b) they comply with the requirements of the animal health certificate referred to in point (a) in respect of the territories or third countries that they come from and any territories or third countries they transit, as referred to in paragraphs 1(a), (b) and (c) of Article 1.

Article 3

Repeals

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Article 4

Transitional provisions

For a transitional period until 29 April 2015, Member States shall authorise imports into the Union of dogs, cats or ferrets which are accompanied by a health certificate issued not later than 28 December 2014 in accordance with the models set out in the Annex to Decision 2005/64/EC or in Annex I to Implementing Decision 2011/874/EU.

Article 5

Applicability

This Decision shall apply from 29 December 2014.

Article 6

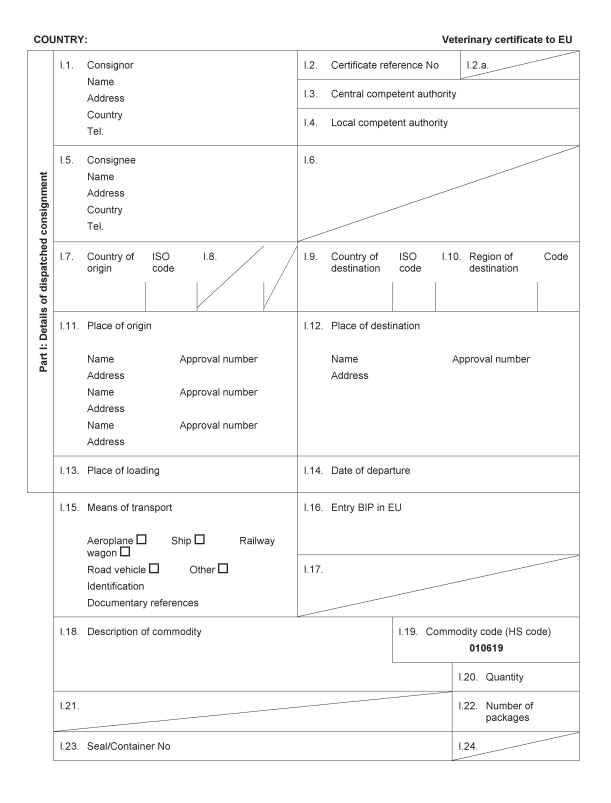
Addressees

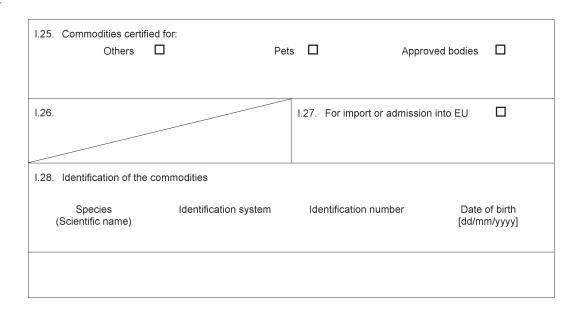
This Decision is addressed to the Member States.

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Model animal health certificate for imports into the Union of dogs, cats and ferrets





						imports	into the Unior	i oi uogo	, outs, icricis
		П.	Health in	formation	II.a. Certif	icate refere	nce No	II.b.	
		-	,	ersigned official vete ountry) certify that the					. (insert name
ion			II.1.	come from holding the competent aut where the animals ensuring the welfa	thority and are are examined	e not subject d regularly a	to any ban on	animal h	ealth grounds,
Part II: Certification			II.2.	showed no signs journey at the tim authority within 48	e of examinat	ion by a vet	erinarian autho		
Part II:		(¹) either	[11.3.	are destined for a body, institute or centre described in Box I.12 and approved in accordance with Annex C to Council Directive 92/65/EEC, and come from a territory or third country listed in Annex II to Commission Implementing Regulation (EU) No 577/2013.]					
(¹) or [II.3. were at least 12 wee 21 days have elaps vaccination (²) carried Annex III to Regulatic Council, and any sub validity of the precedir				elapsed since rried out in ac lation (EU) No subsequent r	e the com cordance wi 576/2013 c evaccination	pletion of th th the validity of the Europear	e primai requireme n Parliam	ry anti-rabies ents set out in ent and of the	
			(¹) either	Commiss	sion Implemer	nting Regula	third country tion (EU) No 5 are provided in	577/2013	and details of
			(¹) or	third cou Part 1 of rabies ar the vete 30 days date of greater t out withi details of	y come from or are scheduled to transit through, a territory of d country listed in Annex I to Commission Decision 2004/211/EC or i t 1 of Annex II to Commission Regulation (EU) No 206/2010, and ies antibody titration test (⁴), carried out on a blood sample taken b veterinarian authorised by the competent authority not less tha days after the preceding vaccination and at least 3 months prior to th e of issue of this certificate, proved an antibody titre equal to c ater than 0,5 IU/ml (⁵) and any subsequent revaccination was carrie within the period of validity of the preceding vaccination, and th ails of the current anti-rabies vaccination and the date of sampling for ting the immune response are provided in the table below:				4/211/EC or in 6/2010, and a mple taken by not less than ths prior to the e equal to or on was carried ation, and the of sampling for
									V:
	1	Fransponder o	or tattoo	_			Validity of vac		v:
	Alp num code anir	ha- heric in of the r	Date of nplantation and/or eading (°) d/mm/yyyy]	Date of vaccination [dd/mm/yyyy]	Name and manufacturer of vaccine	Batch number			v: Date of blood sampling [dd/mm/yyyy]
	Alp num code	ha- heric in of the r	Date of nplantation and/or eading (⁶)	vaccination	manufacturer	Batch	Validity of vac	cination	Date of blood
	Alp num code	ha- heric in of the r	Date of nplantation and/or eading (⁶)	vaccination	manufacturer	Batch	Validity of vac	cination	Date of blood
	Alp num code	ha- heric in of the r	Date of nplantation and/or eading (⁶)	[II.4. are dogs Delegate Echinoco by the Commiss	s destined for ccus multiloc administering	Batch number a Member 3 (EU) No 115 <i>ularis</i> , and t veterinaria ed Regulatic	Validity of vac	Annex I to ve been t e treatmence with	Date of blood sampling [dd/mm/yyyy]

COUNTRY

Imports into the Union of dogs, cats, ferrets

	-		-		
п.	Health info	rmation	II.a. Certificate referen	nce No	II.b.
Transp	onder or tattoo	Anti-echinoc	occus treatment	Administerii	ng veterinarian
	neric code of the dog	e Name and manufacturer of the product	manufacturer of the time of treatment Name in capitals, sta		
	Notes				
	(a)	This certificate is meant ferrets (<i>Mustela putorius</i>	for dogs (Canis lupus fam furo).	iliaris), cats (Felis .	silvestris catus) ar
	(b)	the case of transport by	or 10 days from the date sea, that period of 10 da ation of the journey by sea	ys is extended by a	
	Part I:				
	Box I.11:	<i>Place of origin</i> : name a registration number.	nd address of the dispate	ch establishment. I	ndicate approval
	Box I.12:		ndatory where the anima dance with Annex C to Co		
	Box I.25:		<i>r</i> : indicate 'others' where lation (EU) No 576/2013 (
	Box I.28:	Identification system: sel	ect transponder or tattoo.		
		Identification number. ind	dicate the transponder or t	attoo alphanumerio	code.
	Part II:				
	(1)	Keep as appropriate.			
	(2)		be considered a primary ity of a previous vaccination		vas not carried o
	(3)	A certified copy of the is shall be attached to the c	identification and vaccina certificate.	tion details of the	animals concerne
	(4)	The rabies antibody titra	tion test referred to in poin	t II.3.1:	
		competent authorit	ut on a sample collected sy, at least 30 days after		
		before the date of i	mport;		
			evel of neutralising antiboo	dy to rabies virus i	

 does not have to be renewed on an animal, which following that test wis satisfactory results, has been revaccinated against rabies within the period validity of a previous vaccination. A certified copy of the official report from the approved laboratory on the result of thrabies antibody test referred to in point II.3.1 shall be attached to the certificate. By certifying this result, the official veterinarian confirms that he has verified, to the be 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 te referred to in point II.3.1. In conjunction with footnote (³), the marking of the animals concerned by the implantation of a transponder or by a clearly readable tattoo applied before 3 July 20° must be verified before any entry is made in this certificate and must always precede an vaccination, or where applicable, testing carried out on those animals. The treatment against <i>Echinococcus multilocularis</i> referred to in point II.4 must: be administered by a veterinarian within a period of not more than 120 hours ar not less than 24 hours before the time of the scheduled entry of the dogs into or of the Member States or parts thereof listed in Annex I to Commission Delegate Regulation (EU) No 1152/2011; consist of an approved medicinal product which contains the appropriate dose praziquantel or pharmacologically active substances, which alone or combination, have been proven to reduce the burden of mature and immatu intestinal forms of <i>Echinococcus multilocularis</i> in the host species concerned. The table referred to in point II.4 must be used to document the details of a furth 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	L				0	II.b.
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Date: Signature:		Official ve	terinarian			
		Nam	ne (in capital letters):		Qualifica	tion and title:
Stamp:		Date	9:		Signatur	e:
		Star	np:			

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PART 2

Explanatory notes for completing the animal health certificates

- (a) Where the 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 official veterinarian, or completely deleted from the certificate.
- (b) 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.
- (c) The 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(s) of another Member State, and accompanied, if necessary, by an official translation.

- (d) If for reasons of identification of the items of the consignment (schedule in point I.28 of the model animal health certificate), additional sheets of paper or supporting documents are attached to the certificate, those sheets of paper or documents shall also be considered as forming part of the original of the certificate by the application of the signature and stamp of the official veterinarian, on each of the pages.
- (e) When the certificate, including additional sheets or documents referred to in point (d), 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.
- (f) The original of the certificate shall be completed and signed by an official veterinarian of the exporting territory or third country. The competent authority of the exporting territory or third country shall ensure that rules and principles of certification equivalent to those laid down in Council Directive 96/93/EC 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 water-marked.

(g) The certificate reference number referred to in Boxes I.2 and II.a shall be issued by the competent authority of the exporting territory or third country.

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