ANNEX

PART 1

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

co	UNTR	łY:	8 .5		Veterinary certificate to EU	
	I.1.	Consignor	I.2. Certificate refer	ence No	1.2.a.	
		Name Address	I.3. Central compet	ent authority	•	
		Country Tel.	I.4. Local competer	nt authority		
	1.5.	Consignee Name	1.6.			
nen		Address				
ignı		Country		<u> </u>	e	
d cons		Tel.			8	
patche	1.7.	Country ISO code I.8. of origin	I.9. Country destination	of ISO code	I.10. Region of Code destination	
Part I: Details of dispatched consignment						
Detai	I.11. Place of origin		I.12. Place of destina	tion		
art I:		None	N		A	
ď		Name Approval number Address	Name Address	/	Approval number	
		Name Approval number	Address			
0		Address	· · · ·			
		Name Approval number				
8		Address	а 2 ⁶ ал			
	1.13.	Place of loading	I.14. Date of departur	e		
					r	
2	l.15.	Means of transport	I.16. Entry BIP in EU		ga 2 D	
24					2	
		Aeroplane Ship Railway wagon		a 8	·	
	Road vehicle		1.17.			
ŝ						
	l.18.	Description of commodity		I.19. Commo	dity code (HS code) 010619	
		· · · · · · · · · · · · · · · · · · ·			I.20. Quantity	
	1.21.				I.22. Number of packages	
	1.23.	Seal/Container No	5.	~	1.24.	



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	II. I	lealth inf	formation		II.a.	Certificate refe	erence No	ll.k)
			fficial veterinaria n Box I.28:	an of		(ins	ert name of th	ird country) certify that t
		1Í.1.	authority an	holdings or busin d are not subject t d which comply with	o any ban on	animal health	grounds, where	the anima	als are examine
		II.2.		signs of diseases by a veterinarian a					
	(¹) either	[11.3,	Annex C to (l for a body, institu Council Directive 9: Implementing Reg	2/65/EEC, and	come from a te			
	(1) or	[1].3.	since the con requirements the Council,	t 12 weeks old at the npletion of the print s set out in Annex and any subsequ ccination (³), and	nary anti-rabies III to Regulation	vaccination (²) on (EU) No 576) carried out in a 5/2013 of the E	accordance uropean P	e with the validi arliament and o
		(¹) either	in Annex II to	om, and in case of Commission Imp ation are provided	lementing Reg	ulation (EU) No	577/2013 and		
	* * * *	(1) or	Annex II to	rom or are sched Commission Regu Implementing Regu	ulation (EU) N	o 206/2010 o	rritory or third or listed without	country list time limi	ted in Part 1 o t in Annex I t
×		*	authoris least thr greater validity	antibody titration ed by the compete ee months prior to than 0,5 IU/ml (⁵) a of the preceding v ided in column 8 in	nt authority not the date of iss and any subse accination, and	less than 30 d ue of this certin quent revaccin the date of s	ays after the pro ficate, proved a ation was carrie	eceding va n antibody ed out with	titre equal to o titre equal to o tin the period o
	Tra	nsponder o	r tattoo	10 N.	1	1.			
						2	Validity of vac	cination	×
	Alphanum code of the a	inimal ai	Date of implantation nd/or reading (⁶) [dd/mm/yyyy]	Date of vaccination [dd/mm/yyyy]	Name and manufacturer o vaccine	f Batch number	From [dd/mm/yyyy]	to [dd/mm/yyyy]	Date of blood sampling [dd/mm/yyyy]
	1		.2	3	4	5	6	7	8
		Ð			2				
	*		a a					× ×	3
					5 (A.				
8			<i>d</i> = 2		,		the second s		
(¹) either	[1].4.		ent includes dogs Regulation (EU)					

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COUN	ITRY		8 	Impor	ts into the Unio	n of dogs, cats, ferrets	
11.	Health information	88	II.a.	Certificate refere	ince No	II.b.	
	Transponder or tattoo			atment	Administ	Administering veterinarian	
	anumeric code of the dog	Name and manufacturer of the product			Name in capitals, stamp and signature		
7	2000 - 20000 - 2000 - 2000 - 2000 - 20000 - 2000 - 2000 - 2000 - 2000 -	an a			a 24 S		
2	9 9			8	а. 	· · · ·	
	n a chairte an tha a can tha can a tha an	· ·	-				
8							
	×	N		n	were notediation war in	5 	
н. ²²		5 6) 1	-1	
(1) or	[II.4. the dogs	forming part of the consign	nment h	ave not been treated	against Echinoc	occus multilocularis.]	
Notas				ne na seconda en esta esta esta esta esta esta esta esta		5	
Notes		× 1			5 c		
This ce of 10 d	ertificate is valid for 10 day ays is extended by an add	rs from the date of issue b litional period correspondi	y the off ng to the	ficial veterinarian. In e duration of the journ	the case of trans ney by sea.	port by sea, that period	
	i K						
Part I:				а – с. 		. *	
Box I.1	1: Place of origin: name	and address of the dispa	tch esta	blishment. Indicate a	pproval or registr	ation number.	
Box I.1		: mandatory where the a					
Dort I. I		ex C to Council Directive S			body, montato		
Box I.2	5: Commodities certified	d for: indicate			ž		
		is (Canis lupus familiaris) ance with Article 5(4) of R					
		s' where dogs, cats or fer approved body, institute of					
	- 'others' where do	gs, cats or ferrets are mov	red in ac	cordance with Article	10 of Council D	irective 92/65/EEC.	
Box I.2	8: Identification system:	select transponder or tatte	00.	2 ^{- 18}			
	Identification number.	indicate the transponder	or tattoo	alphanumeric code.		5.	
Part II:	*			194 194	л». К		
(¹) Kee	p as appropriate.						
	revaccination must be c vious vaccination.	considered a primary vac	cination	if it was not carried	d out within the	period of validity of a	
(³) A c	ertified copy of the identific	cation and vaccination deta	ails of th	e animals concerned	shall be attache	d to the certificate.	
(⁴) The	rabies antibody titration te	est referred to in point II.3:		з У.,	الغري	× •	
		sample collected by a ve on and three months befo			competent auth	oority, at least 30 days	

must measure a level of neutralising antibody to rabies virus in serum equal to or greater than 0,5 IU/ml;

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11.	Health information	II.a.	Certificate reference No	II.b.
11.	Health Information	II.a.	Certificate reference No	11.D.
	must be performed by a laboratory approved in approved laboratories available at http://ec.europ			
	does not have to be renewed on an animal, which against rables within the period of validity of a pre-			nas been revaccinated
	certified copy of the official report from the approv int II.3 shall be attached to the certificate.	ed labora	tory on the result of the rabies antib	ody test referred to ir
wit	certifying this result, the official veterinarian confirm h contacts with the laboratory indicated in the re tibody titration test referred to in point II.3.	ns that he port, the	has verified, to the best of his ability authenticity of the laboratory report	and where necessary on the results of the
rea	conjunction with footnote (3), the marking of the an adable tattoo applied before 3 July 2011 must be v cede any vaccination, or where applicable, testing of	verified be	efore any entry is made in this certific	
⁷) The	e treatment against Echinococcus multilocularis refe	erred to in	point II.4 must:	ge K I
_	be administered by a veterinarian within a period time of the scheduled entry of the dogs into o	ne of the		
	Commission Implementing Regulation (EU) 2018/	878;		
	consist of an approved medicinal product which active substances, which alone or in combinatior intestinal forms of <i>Echinococcus multilocularis</i> in th	contains n, have be	een proven to reduce the burden of	or pharmacologically
the	consist of an approved medicinal product which active substances, which alone or in combinatior	contains n, have be he host sp ument the entry int	een proven to reduce the burden of pecies concerned. details of a further treatment if admir	or pharmacologically mature and immature nistered after the date
the Anr	consist of an approved medicinal product which active substances, which alone or in combination intestinal forms of <i>Echinococcus multilocularis</i> in the table referred to in point 'II.4 must be used to docu certificate was signed and prior to the scheduled	contains n, have be he host sp ument the entry int	een proven to reduce the burden of pecies concerned. details of a further treatment if admir	or pharmacologically mature and immature nistered after the date
the Anr Official	consist of an approved medicinal product which active substances, which alone or in combinatior intestinal forms of <i>Echinococcus multilocularis</i> in the table referred to in point II.4 must be used to docu certificate was signed and prior to the scheduled nex to Commission Implementing Regulation (EU) 2	contains n, have be he host sp ument the entry int	een proven to reduce the burden of pecies concerned. details of a further treatment if admir	or pharmacologically mature and immature nistered after the date s thereof listed in the
the Anr Official	consist of an approved medicinal product which active substances, which alone or in combinatior intestinal forms of <i>Echinococcus multilocularis</i> in the table referred to in point II.4 must be used to docu certificate was signed and prior to the scheduled nex to Commission Implementing Regulation (EU) 2 veterinarian me (in capital letters):	contains n, have be he host sp ument the entry int	een proven to reduce the burden of the becker concerned. details of a further treatment if admir o one of the Member States or parts	or pharmacologically mature and immature nistered after the date s thereof listed in the
official Nar Dat	consist of an approved medicinal product which active substances, which alone or in combinatior intestinal forms of <i>Echinococcus multilocularis</i> in the table referred to in point II.4 must be used to docu certificate was signed and prior to the scheduled nex to Commission Implementing Regulation (EU) 2 veterinarian me (in capital letters):	contains n, have be he host sp ument the entry int	een proven to reduce the burden of the becies concerned. details of a further treatment if admir o one of the Member States or parts Qualification and title	or pharmacologically mature and immature nistered after the date s thereof listed in the
fficial Nar	consist of an approved medicinal product which active substances, which alone or in combinatior intestinal forms of <i>Echinococcus multilocularis</i> in the table referred to in point II.4 must be used to doct certificate was signed and prior to the scheduled nex to Commission Implementing Regulation (EU) 2 veterinarian me (in capital letters):	contains n, have be he host sp ument the entry int	een proven to reduce the burden of the becies concerned. details of a further treatment if admir o one of the Member States or parts Qualification and title	or pharmacologically mature and immature nistered after the date s thereof listed in the
fficial Nar	consist of an approved medicinal product which active substances, which alone or in combinatior intestinal forms of <i>Echinococcus multilocularis</i> in the table referred to in point II.4 must be used to doct certificate was signed and prior to the scheduled nex to Commission Implementing Regulation (EU) 2 veterinarian me (in capital letters):	contains n, have be he host sp ument the entry int	een proven to reduce the burden of the becies concerned. details of a further treatment if admir o one of the Member States or parts Qualification and title	or pharmacologically mature and immature nistered after the date s thereof listed in the

Explanatory notes for completing the animal health certificate

- Where the certificate states that certain statements shall be kept as appropriate, statements which are not relevant (a) 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.
- The certificate shall be drawn up in at least one of the official languages of the Member State of the border (c) 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.
- If for reasons of identification of the items of the consignment (schedule in point I.28 of the model animal health (d) 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.

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- (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 Directive 96/93/EC are followed.
- (g) 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.
- (h) 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.