Source: Regulation(EU) Nr. 2019/1293 Annex IV

Part I

Model health certificate for the non-commercial movement of dogs, cats or ferrets to a member state from a third country or region according to article 5, section 1 and 2, of Regulation (EU) No. 576/2013

COU	VTRY					Veterinary certificate to EU
	I.1.	Consignor	1.2.	Certificate	e reference No	I.2.a.
		Name Address	1.3.	Central co	ompetent auth	prity
			1.4.	Local com	npetent author	ity
t		Tel.				
gnme	I.5.	Consignee	I.6.	Operator in the EU		r the consignment
onsi		Name				
shed c		Address				
spato		Postal code				
of di		Tel.				
Part I: Details of dispatched consignment	1.7.	Country of origin ISO I.8. Region of Code origin	1.9.	Country o destinatiø	f ISC n code	
	I.11	Place of origin	I.12.	Place of c	destination	
	I.13.	Place of loading	I.14.	Date of de	eparture	
	I.15.	Means of transport	I.16.	Entry BIP	in EU	
			I.17.	No.(s) of	CITES	
					1.40.0	111
	1.18.	Description of commodity			1.19. Commo	odity code (HS code) 010619
				,		010013
	-				1.20	). Quantity
	I.21.	Temperature of products			1.22	2. Total number of packages
	1.23.	Seal/Container No			1.24	I. Type of packaging

cour	NTRY:							Veterinary certificate to EU
	I.25. Commoditie Pets	s certifie	d for:					
	I.26. For transit to	o 3 <sup>rd</sup> Cou	ntry			I.27. For ir	mport or admission into E	U
	I.28. Identification	n of the c	ommoditie	3				
	Species (Scientific name)	Sex	Colour	Breed	Identification nu	mber	Identification system	Date of birth [dd/mm/yyyy]

## 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

COUNTRY							Article 5(1) and (	2) of Regulation	n (EU) No 5	76/20	)13
II.	Health	n inf	formation			II.a. Certificate	e reference No	II.b.			
					veterinarian ( <i>ins</i> e		authorised by tl		authority	(1)	of
	Purpo	se/	nature of jo	urney attest	ted by the owne	er:					
	II.1.		to carry ou states that authorisati the owner	It the non-c t the anima on in writing within not or a trans	commercial mov als described i g from the owne more than five	vement of the an n Box I.28 will er to carry out th days of his mov	al person who has au imals on behalf of the accompany the own e non-commercial mo rement and are not s the non-commercial	e owner, suppor ler or the natur ovement of the a subject to a mov	ted by evide al person v animals on b vement that	ence vho h behall aims	( <sup>3</sup> ), nas f of s at
( <sup>1</sup> ) e	either		[the owner	;]							
( <sup>1</sup> ) c	or		10-00		no has authorisa alf of the owner		om the owner to carry	y out the non-cor	mmercial mo	ovem	ent
(1) (	or				designated by nals on behalf o		acted by the owner	to carry out t	he non-con	nmero	cial
(1) ei	ther [l	1.2.	the animal	s described	l in Box I.28 are	moved in a num	ber of five or less;]				
( <sup>1</sup> ) or	. [1	1.2.	are going	to participat	te in competitio	ns, exhibitions o	nber of more than five r sporting events or i s provided evidence	in training for the	ose events,	and	the
( <sup>1</sup> ) e	either		[to attend s	such event;]	]						
( <sup>1</sup> ) c	or		[with an as	sociation or	rganising such	events;]					
	Attest	atio	n of rabies	vaccination	and rabies ant	ibody titration tes	: <u>t:</u>				
( <sup>1</sup> ) ei	ther [l	1.3.	vaccination least have	n, or are bei e not elaps	tween 12 and 1 sed since the	6 weeks old and completion of	12 weeks old and have received an ar the primary vaccina nnex III to Regulation	nti-rabies vaccina ition against ral	ation, but 21 bies carried	days	s at
			II.3.1	Implement I.5 has info	ing Regulation	(EU) No 577/20	of the animals indica 13 and the Member S ses the movement of	State of destinati	ion indicated	d in E	Box
( <sup>1</sup> ) eith	er		[11.3.2	from birth	until the time of		or the natural person ercial movement the es;]				
( <sup>1</sup> ) or			[11.3.2	their birth a		accination which	nd it can be establis complied with the va				
( <sup>1</sup> ) or	/and [l	1.3.	least 21 d accordanc	lays have e e with the	elapsed since t validity require	the completion of ments set out i	eeks old at the time of the primary anti-ra n Annex III to Regu eriod of validity of the	abies vaccinatio ulation (EU) No	n ( <sup>4</sup> ) carrie 576/2013 a	d out and a	in
( <sup>1</sup> ) eith	er		[II.3.1	Implement listed in Au country ot accordance	ing Regulation nnex II to Impl her than those with point (c)	(EU) No 577/20 ementing Regula e listed in Anne of Article 12(1) of	from a territory or a 113, either directly, th ation (EU) No 577/20 x II to Implementing f Regulation (EU) No d in the table below;]	nrough a territor 013 or through a 9 Regulation (El 0 576/2013 ( <sup>7</sup> ), a	y or a third a territory o U) No 577/	cour r a th 2013	ntry nird in

## Non-commercial movement into a Member State from a territory or third country of dogs, cats or ferrets in accordance with

II. Health information						Certificato	8. 12		-	(EU) No 576/2013	
	Health Ir		120	765. 9. 11. 1557			reference N	1992 1997 - 1997 - 1997 - 1997	II.b.		
( <sup>1</sup> ) or		[II.3.1	count rabie by the prece an at carrie curre	nimals described in try other than those s antibody titration t e competent author eding vaccination ar ntibody titre equal ed out within the p nt anti-rabies vacc ded in the table belo	e listed est ( <sup>8</sup> ), ity on th d at lea to or gi eriod of nation	in Annex I carried out le date ind st three m reater than validity o	I to Implem on a blood icated in the onths prior 0 0,5 IU/ml f the prece	enting Regu sample take table below to the date o ( <sup>9</sup> ) and any ding vaccina	lation (EU) Normality of the veter of the veter of the veter of the veter of the veter of the veter of the subsequent of the veter of t	lo 577/2013 and a rinarian authorised n 30 days after the certificate, proved revaccination was the details of the	
Tr	anspond	der or tatto	D						dity of ination		
code	numeric of the mal [dd/mm/yyy		tation vaccination /or [dd/mm/yyyy] g ( <sup>10</sup> )		Name and manufacturer of vaccine		Batch number	From [dd/mm/yyyy]	to [dd/mm/yyyy]	Date of the blood sampling [dd/mm/yyyy]	
				-							
								r			
				1			1	1	1	]	
Implementing Re multilocularis, and accordance with				ogs described in Bo ementing Regulation ocularis, and the co	scribed in Box I.28 are destined for a Member State listed in the Annex to Commis g Regulation (EU) 2018/878 and have been treated against Echinoco s, and the details of the treatment carried out by the administering veterinaria with Article 6 of Commission Delegated Regulation (EU) 2018/772 ( <sup>11</sup> ) ( <sup>12</sup> ) ( <sup>13</sup> )						
(1) or		[1].4.	the d	ogs described in Bo	- x I.28 h	ave not be	en treated a	igainst <i>Echin</i>	ococcus muli	ilocularis ( <sup>11</sup> ).]	
				Anti-	echinoo	occus tre	atment		Administe	ring veterinarian	
		r or tattoo the dog	٩	Name and manufac of the product			d/mm/yyyy] reatment [0			capitals, stamp signature	

cou	NTRY		٢	or third country of dogs, o	o a Member State from a territory cats or ferrets in accordance with 2) of Regulation (EU) No 576/2013
II.	He	alth information	II.a.	Certificate reference No	II.b.
Note	es		10		
(a)		This certificate is meant for dogs ( <i>Canis lu furo</i> ).	pus t	familiaris), cats (Felis silvestris c	catus) and ferrets (Mustela putorius
(b)		This certificate is valid for 10 days from documentary and identity checks at http://ec.europa.eu/food/animal/liveanimals/	the	designated Union travellers	
		In the case of transport by sea, that perio duration of the journey by sea.	d of	10 days is extended by an add	itional period corresponding to the
		For the purpose of further movement int documentary and identity checks for a total vaccination or until the conditions relating to whichever date is earlier. Please note that territory of animals less than 16 weeks old http://ec.europa.eu/food/animal/liveanimals/	our months or until the date of ex mals less than 16 weeks old refe rtain Member States have infor erred to in point II.3 is not autho	piry of the validity of the anti-rables erred to in point II.3 cease to apply, med that the movement into their	
Parl	t I:				
Box	1.5:	Consignee: indicate Member State of first d	estin	ation.	
Box	1.28:	Identification system: select of the following	: tran	sponder or tattoo.	
		Identification number. indicate the transpon	der o	r tattoo alphanumeric code.	
		Date of birth/breed: as stated by the owner.			
Parl	: 11:				
(1)	Keep	as appropriate.			
(2)		leclaration referred to in point II.1 shall be ements set out in Part 3 of Annex IV to Imple			
( <sup>3</sup> )	proof	vidence referred to in point II.1 (e.g. boardir of membership) shall be surrendered on req nt (b) of the Notes.			
(4)		evaccination must be considered a primary ous vaccination.	vac	cination if it was not carried ou	t within the period of validity of a
( <sup>5</sup> )		eclaration referred to in point II.3.2 to be at ements laid down in Parts 1 and 3 of Annex I			
(6)	A cert	ified copy of the identification and vaccination	n deta	ails of the animals concerned sha	all be attached to the certificate.
(7)	by the had n perim to Im	nird option is subject to the condition that the e competent authorities responsible for the cl o contact with animals of species suscepti eter of an international airport during the tran plementing Regulation (EU) No 577/2013. ements set out in Parts 2 and 3 of Annex I to	hecks ble c sit thi This	s referred to in point (b), a declar of rabies and remain secure with rough a territory or a third countr declaration shall comply with	ration stating that the animals have hin the means of transport or the y other than those listed in Annex I the format, layout and language

II.       Health information       II.a. Certificate reference No       II.b.         (*)       The rables 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 rables 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 rables 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 rables antibody test referred to in point II.3.1.         (*)       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 rables any vaccination, or where applicable, testing carried out on the antimatic.         (*)       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 thos animals.         (*)	cour	NTRY	I	or third country of dogs, o	o a Member State from a territory cats or ferrets in accordance with 2) of Regulation (EU) No 576/2013		
<ul> <li>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;</li> <li>must measure a level of neutralising antibody to rabies virus in serum equal to or greater than 0,5 IU/ml;</li> <li>must be performed by a laboratory approved in accordance with Article 5 of Council Decision 2000/258/EC (list of approved laboratories available at http://ec.europa.eu/food/animal/liveanimals/pets/approval_en.htm);</li> <li>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.</li> <li>A certified copy of the official report from the approved laboratory on the results of the rabies antibody test referred to in point III.3.1.</li> <li>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.</li> <li>In conjunction with footnote (6), the marking of the animals concerned by the implantation of a transponder or by a clearly recede any vaccination, or where applicable, testing carried out on those animals.</li> <li>The treatment against <i>Echinococcus multilocularis</i> referred to in point II.4 must:</li> <li>be administered by a veterinarian within period of no 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 the Annex to Implementing Regulation (EU) 2018/878.</li> <li>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 an</li></ul>	II.	Health information	II.a.	Certificate reference No	II.b.		
<ul> <li>the date of vaccination and three months before the date of import;</li> <li>must measure a level of neutralising antibody to rabies virus in serum equal to or greater than 0.5 IU/ml;</li> <li>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);</li> <li>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.</li> <li>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.</li> <li>(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.</li> <li>(10) 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 3.1uly 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.</li> <li>(11) The treatment against <i>Echinococcus multilocularis</i> referred to in point II.4 must:</li> <li>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 sobstance of the dogs into one of the Member States or parts thereof listed in the Annex to Implementing Regulation (EU) 2018/878.</li> <li>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 ma</li></ul>	(8)	The rabies antibody titration test referred to in poin	t II.3.	1:			
<ul> <li>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);</li> <li>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 11.3 ishall be attached to the certificate.</li> <li>By certifying this result, the official report from the approved laboratory on the results of the rabies antibody test referred to in point 11.3.1.</li> <li>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 11.3.1.</li> <li>In conjunction with footnote (B), the marking of the animals concerned by the implantation of a transponder or by a clearly readable tatoo 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.</li> <li>The treatment against <i>Echinococcus multilocularis</i> referred to in point 11.4 must:</li> <li>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 substances, which alone or in combination, have been proven to reduce the burden of mature and immature intestinal forms of <i>Echinococcus multilocularis</i> in the host species concerned.</li> <li>The table referred to in point 11.4 must be used to document the details of reatments if administered after the date the certificate was signed and prior to the scheduled entry into</li></ul>					ent authority, at least 30 days after		
laboratories available at http://ec.europa.eu/food/animal/liveanimals/pets/approval_en.htm);		must measure a level of neutralising antibody to ra	bies	virus in serum equal to or greate	r than 0,5 IU/ml;		
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 11.3.1 shall be attached to the certificate.         (*)       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 11.3.1.         (19)       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.         (11)       The treatment against Echinococcus multilocularis referred to in point 11.4 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 the Annex to 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.         (12)       The table referred to in point 11.4 must be used to document the details of a further treatment if administered after the date the certificate was signed and							
<ul> <li>point II.3.1 shall be attached to the certificate.</li> <li>(*) 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.</li> <li>(*) 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.</li> <li>(*) The treatment against <i>Echinococcus multilocularis</i> referred to in point II.4 must:</li> <li>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 the Annex to Implementing Regulation (EU) 2018/878;</li> <li>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 <i>Echinococcus multilocularis</i> in the host species concerned.</li> <li>(**) 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 point (b) of the Notes and Annex to Implementing Regulation (EU) 2018/878.</li> <li>(**) 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 described in point (b) of the Notes and in conjunction with footnote (11).&lt;</li></ul>	-				ory results, has been revaccinated		
<ul> <li>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.</li> <li>(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 very is made in this certificate and must always precede any vaccination, or where applicable, testing carried out on those animals.</li> <li>(11) The treatment against <i>Echinococcus multilocularis</i> referred to in point II.4 must: <ul> <li>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 the Annex to Implementing Regulation (EU) 2018/878;</li> <li>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 <i>Echinococcus multilocularis</i> in the host species concerned.</li> <li>(1°) 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 the Annex to Implementing Regulation (EU) 2018/878.</li> <li>(1°) 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).</li> </ul> Official veterinarian/Authorised veterinarian Name (in capital letters): <ul> <li>Qualification and title:</li> <li>Address</li> <li>Telephone:</li> <li>Date:</li> <li>Signature:</li> </ul></li></ul>			rove	d laboratory on the results of the	e rabies antibody test referred to in		
<ul> <li>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.</li> <li>(1) The treatment against <i>Echinococcus multilocularis</i> referred to in point II.4 must:         <ul> <li>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 the Annex to Implementing Regulation (EU) 2018/878;</li> <li>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 <i>Echinococcus multilocularis</i> in the host species concerned.</li> <li>(12) 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 the Annex to Implementing Regulation (EU) 2018/878.</li> <li>(13) 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).</li> </ul> </li> <li>Official veterinarian/Authorised veterinarian         <ul> <li>Name (in capital letters): Qualification and title:</li> <li>Address</li> <li>Telephone:</li> <li>Date: Signature:</li> </ul> </li> </ul>	( <sup>9</sup> )	with contacts with the laboratory indicated in the					
<ul> <li>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 the Annex to Implementing Regulation (EU) 2018/878;</li> <li>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 <i>Echinococcus multilocularis</i> in the host species concerned.</li> <li>(12) 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 the Annex to Implementing Regulation (EU) 2018/878.</li> <li>(13) 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).</li> <li>Official veterinarian/Authorised veterinarian</li> <li>Name (in capital letters): Qualification and title:</li> <li>Address</li> <li>Telephone:</li> <li>Date: Signature:</li> </ul>	(10)	readable tattoo applied before 3 July 2011 must	be ve	erified before any entry is made			
<ul> <li>the scheduled entry of the dogs into one of the Member States or parts thereof listed in the Annex to Implementing Regulation (EU) 2018/878;</li> <li>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 <i>Echinococcus multilocularis</i> in the host species concerned.</li> <li>(12) 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 the Annex to Implementing Regulation (EU) 2018/878.</li> <li>(13) 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).</li> <li>Official veterinarian/Authorised veterinarian</li> <li>Name (in capital letters): Qualification and title:</li> <li>Address</li> <li>Telephone:</li> <li>Date: Signature:</li> </ul>	(11)	The treatment against Echinococcus multilocularis	refer	red to in point II.4 must:			
substances, which alone or in combination, have been proven to reduce the burden of mature and immature intestinal forms of <i>Echinococcus multilocularis</i> in the host species concerned. (12) 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 the Annex to Implementing Regulation (EU) 2018/878. (13) 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): Address Telephone: Date: Signature:		the scheduled entry of the dogs into one of the					
the certificate was signed and prior to the scheduled entry into one of the Member States or parts thereof listed in the Annex to Implementing Regulation (EU) 2018/878. ( <sup>13</sup> ) 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): Address Telephone: Date: Signature:		substances, which alone or in combination, have been proven to reduce the burden of mature and immature intestina					
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):         Address         Telephone:         Date:       Signature:	(12)	the certificate was signed and prior to the sched	uled				
Name (in capital letters):Qualification and title:Address	(13)	certificate was signed for the purpose of further mo					
Address Telephone: Date: Signature:	Offic	sial veterinarian/Authorised veterinarian					
Telephone: Date: Signature:		Name (in capital letters):		Qualification and	title:		
Date: Signature:		Address					
		Telephone:					
Stamp:		Date:		Signature:			
		Stamp:					

OUNTRY	Δ	Article 5(1) and (2) of Regulation (EU) No 576/20						
I. Health information	II.a. Certificate refe	erence No	II.b.					
Endorsement by the competent at	uthority (not necessary when the certificate	e is signed by a	an official veterinarian)					
Name (in capital letters):		Qualification a	nd title:					
Address								
Telephone:								
Date:		Signature:						
Stamp:								
Official at the travellers' point of er	ntry (for the purpose of further movement	into other Merr	nber States)					
Name (in capital letters):		Title:						
Address								
Telephone:								
Email address:								
Date of completion of the do	cumentary and identity checks:	Signature:	Stamp:					

## 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