**Export Health Certificate** 

	I.1. Consignor					I.2. IMSOC Ref	Terence				
	Name					I.2.a. Local Reference					
	Address										
	Country ISO Code										
	· · ·					700 1					
	I.S. Consignee						mpetent authority				
Part I : Details of consignment	Name					1.4. Local com	petent authority				
ne	Address			ICO C1-							
Ħ	Country			ISO Code							
Sig	I.7. Country of orig	gin			ISO Code	I.9. Country of	destination			ISO Code	
ol	, ,										
Ę	I.8. Region of origin	n			Code	I.10. Region of	doctination				
S	I.11. Place of Dispa				Code	I.12. Place of d					
ai I	_	itteri				Name	icotification				
et	Name Address					Address					
$\Box$	Approval Number	•				Approval Nui	mhor				
t I	Country	-		ISO Code		Country	iibei		ISO Code		
ar						oo aana y					
4	I.13. Place of Loadi	ing				I.14. Date and	time of departure				
	Name										
	Address										
	Approval Number	•									
	Country			ISO Code							
	145 M					I 4 C P . I . P . I					
	I.15. Means of Tran	_				I.16 Entry Poi	nt				
	Mode	Internation transport	ıal	Identification							
		document									
	I.18. Transport con	nditions					nying documents				
	Ambient $\square$					Commercial document Date of issue					
						reference					
						Country		Place issue	of		
	I.19. Container No	/ Coal No						133410			
	1.19. Container No	/ Seal No									
l	I.20. Certified as					Other 🗆					
	Approved Bodies [			Pets 🗆							
					I 22. For transit through Member State(s)						
- 1	I.21. For transit through a third country					I.22. For transit through Member State(s)					
- 1	Country			ISO Code							
	EU Exit Authority			BCP code		Country		ISO Co	ode		
	EU Entry			BCP code							
	Authority					TOP 75-4-1	no rusiaht				
	I.24. Total quantity  I.28. Description of consignment						I.25. Total gross weight				
1. 01 LIVE ANIMALS											
	0106 Other live										
	Mammals:										
	<b>010619</b> Oth	ner									
	0106190	<b>0</b> Other									
	Commodity		Specie	es	Identification	svstem	Identification number	er	Quantity		
Date of Birth											
	_ att of birth										
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# **EUROPEAN UNION**

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I	EU	ROPEAN U	JNION		(GB) Dogs, cats and ferrets from EU countries / GBHC157E (v3.1)						
		II. Health info	rmation								
		I, the unde described i	rsigned offi n Box I.28:	cial veterir	narian of (insert name of third country) certify that the animals						
		II.1.	come from holdings or businesses described in Box 1.11 which are registered by the competent authority and are not subject to any ban on animal health grounds, where the animals are exam regularly and which comply with the requirements ensuring the welfare of the animals held;								
Part II: Certification	tication	II.2.	examinati	showed no signs of diseases and were fit to be transported for the intended journey at the time of examination by a veterinarian authorised by the competent authority within 48 hours prior to the time of dispatch;							
	art II: Cert	(1)	○ either	[II.3.	are destined for a body, institute or centre described in Box 1.12 and approved in accordance with Annex C to Council Directive 92/65/EEC, and come from a territory or third country listed in Annex 2 to Commission Implementing Regulation (EU) No 577/2013.]						
		(1) or	∘ or	[II.3.	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 (2) carried out in accordance with the validity requirements set out in Annex 3 to Regulation (EU) No 576/2013, and any subsequent revaccination was carried out within the period of validity of the preceding vaccination (3), and						
		(1)	∘ either	country li	ne from, and in case of transit are scheduled to transit through, a territory or third listed in Annex 2 to Commission Implementing Regulation (EU) No 577/2013 and f the current anti-rabies vaccination are provided in columns 1 to 7 in the table						
		(1)(9)(10)	or	document Commissi to 'equida	ne from or are scheduled to transit through, a territory or third country listed in a at relating to 'fresh meat of ungulates' published on gov.uk, in accordance with sion Regulation (EU) No 206/2010 or listed without time limit in a document relating ae' published on gov.uk, in accordance with Commission Implementing Regulation 8/659, and						
			-	details of below, an	f the current anti-rabies vaccination are provided in columns 1 to 7 in the table and						
			-	authorise and at lea titre equa within the	antibody titration test (4), carried out on a blood sample taken by the veterinarian ed by the competent authority not less than 30 days after the preceding vaccination ast three months prior to the date of issue of this certificate, proved an antibody al to or greater than 0,5 IU/ml (5) and any subsequent revaccination was carried out the period of validity of the preceding vaccination, and the date of sampling for the immune response are provided in column 8 in the table below:]						

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Ε	UROPEAN U	JNION		(	(GB) Dogs,	cats and fe	errets from	EU count	ries / GBHC157E (v3.1)
	II. Health info	rmation							
		Transpon der or tattoo					Validity of	Vaccinatio	on
201700 111100		Alphanum eric code of the animal			Name and manufact urer of vaccine		From [dd/mm/y yyy]	To [dd/mm/y yyy]	Date of blood sampling [dd/mm/yyyy]
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		1	2	3	4	5	6	7	8

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Transpon Anti- der or Echinococ tattoo cus alphanum treatment eric code of the dog  Name Date and (add/mm/y manufact yyy) and urer of the time of product treatment [00:00]  Name in capitals, stamp and signature [00:00]  Name in capitals, stamp and signature [00:00]	E	UROPEAN UNION		(	(GB) Dogs, cats and ferrets from EU co	ountries / GBHC157E (v3.1)
der or Echinococ tattoo cus alphanum treatment eric code		II. Health information				
Name Date and incapitals, stamp and signature and ure of the time of product treatment [00:00]		der or tattoo alphanum eric code	Echinococ cus		Administering veterinarian	
	Dart II: Cortification	eric code	Name and manufact urer of the product	[dd/mm/y yyy] and time of treatment	Name in capitals, stamp and signature	

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#### **EUROPEAN UNION**

# II. Health information

## Notes

References to European Union legislation within this certificate are references to direct EU legislation which has been retained in Great Britain (retained EU law as defined in the European Union (Withdrawal) Act 2018) and can be viewed on the UK legislation website (legislation.gov.uk).

References to Great Britain in this certificate include Channel Islands and Isle of Man.

This certificate is valid for 10 days from the date of issue by the official veterinarian. 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.

### Part I:

II: Certification

Part I

Box Place of origin: name and address of the dispatch establishment. Indicate approval or reference registration number.

I.11:

Box Place of destination: mandatory where the animals are destined for a body, institute or reference centre approved in accordance with Annex C to Council Directive 92/65/EEC.

I.12:

Box Commodities certified for: indicate

reference I.25:

'Pets' 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;

'Approved bodies' where dogs, cats or ferrets are moved in accordance with Article 13 of Council Directive 92/65/EEC to an approved body, institute or centre as defined in Article 2(c) of that Directive;

'others' where dogs, cats or ferrets are moved in accordance with Article 10 of Council Directive 92/65/EEC.

Box Identification system: select transponder or tattoo.

reference I.28:

Identification number: indicate the transponder or tattoo alphanumeric code.

#### Part II:

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(1) Keep as appropriate.

(2) Any revaccination must be considered a primary vaccination if it was not carried out within the period of validity of a previous vaccination.

(3) A certified copy of the identification and vaccination details of the animals concerned shall be attached to the certificate.

(4) The rabies antibody titration test referred to in point 11.3:

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 than 0,5 IU/ml;

level of neutralisi ng antibody to rabies virus in serum

measure a

serum equal to or greater

- 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/animals/petmovement/approved-labs\_en);

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# **EUROPEAN UNION**

드	JROPEAN (	JNION (GB) Dogs, cats and ierrets from EU countries / GBHC15/E (V3.1)
	II. Health info	rmation
	-	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 result of the rabies antibody
		test referred to in point 11.3 shall be attached to the certificate.
cation	(5)	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.
Part II: Certification	(6)	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 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.
Par	(7)	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 Great Britain.
	-	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.
	(8)	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 Great Britain.
	(9)	A document relating to 'fresh meat of ungulates' for EU and EFTA states published by the Secretary of State, with the consent of the Scottish and Welsh Ministers, may be found here:
		EU and EFTA states approved to export animals and animal products to Great Britain - data.gov.uk
	(10)	A document relating to 'equidae' for EU and EFTA states published by the Secretary of State, with the consent of the Scottish and Welsh Ministers, may be found here:
		TA states approved to export animals and animal products to Great Britain - data.gov.uk
	Certifying Off  Name (in cap  Date of signates)  Stamp	pital letters) Qualification and title

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