Export Health Certificate

	I.1. Versender					I.2. IMSOC-Bez	zugsnumm	ner		
	Name					I.2.a. Lokale Bezugsnummer				
	Adresse									
	Land		ISO-							
	Ländercode									
	I.5. Empfänger					I.3. Zentrale zi	uetändiga	Rehörde		
	Name					I.4. Zuständige				
	Adresse					1. 1. Zuotanaist	or chiefic 1	scriorae		
	Land		ISO-							
	Darra		Länder	code						
ŀ	17 11				100	I O Dantino				100
┪	I.7. Ursprungsland	<u>.</u>			ISO- Ländercode	I.9. Bestimmu	ngsiand			ISO- Ländercode
Tell I										
	I O Hrenningerogie	an an			Code	I 10 Pagion de	ne Poetimn	aun goo nto		
1	I.8. Ursprungsregio	on			Code	I.10. Region de		aungsorts		
						I.12. Bestimm	ungsort			
	Name					Name				
	Adresse					Adresse Zulassungsnummer				
	Zulassungsnumm	er	100			Land ISO-				
	Land		ISO- Län	dercode		Länd 150- Ländercode				
	I.13. Ladeort					I.14. Datum ur	nd Uhrzeit	des Abtransports		
	Name									
	Adresse									
	Zulassungsnumm	er								
	Land		ISO-	dercode						
			Dan	ucreode						
	I.15. Transportmitt	tel				I.16 Entry Point				
	Typ Dokument Identifikation			on						
	I.18. Beförderungs	bedingungen				I.17. Begleitdo	kumente			
	Gefroren □ Gekühlt □ Umgebungstemp Controlled _					Bezugsnum				
	eratur 🗆 🔭 temperature 🗆				Bezugsnum mer des		Ausste	ellungs 1		
						Begleitdoku ments		uatun	ı	
						Land		Ausste	ellungs	
						1		ort		
	I.19. Containernun	nmer/Plombenr	nummer							
ŀ	I.20. Waren zertifiz	ziert für/als								
	Künstliche Vermeh		Breeding [7						
	Runstnene vermer		Diccums L	-						
İ	I.21. Für die Durch	fuhr durch ein	Drittland			I.22. Für die D	urchfuhr (durch Mitgliedstaate	en 🗆	
	Country		ISO-					-		
			Ländercode	e						
	EU Exit Authority					Country ISO- Ländercode				
	EU Entry	TI Entry								
	Authority	authority BCP code								
	I.24. Gesamtmenge	9				I.25. Bruttogesamtgewicht				
ŀ	I.28. Angaben zur v									
	_		_	ANDEDIA	EIT WEDER OF	NIA NINITE NICOTE	INIDECTIE	FEN		
	1. 05 ANDERE WA				re des Kapitels 1 od	lar 3 unganica	har			
	0511 waren tiel 051199 ander	_	igs, ailuerweit t	weuer gen	amm noch mbe	grinien, micht le	meriue 116	re des Mapriers 1/00	ier 5, ungemeß	vai
	051199 ander									
					T.1		7.1		TAT	
	Erzeugnis	Art			Identifikation	snummer	iaentifika	ationskennzeichen	Warenart	
								I		
	Menge Datum der Gewinnung/I				Herstellung		Fertigungsanlage			
Į										

de 1/5

EUROPÄISCHE UNION

EUROPAISCHE UNION (V3											
	II. Gesundhei	sinformatione	n								
Part II: Certification	I, the unde	I, the undersigned, official veterinarian, hereby certify that:									
	II.1.	.1. The exporting country (name of exporting country) (2)									
		II.1.1.	caprine ple	europneumonia and Rift of the ova(1)/embryos(1)	ste des petits ruminants, sheep and goat pox, contagious ft Valley fever during the 12 months immediately prior to to be exported and until their date of dispatch to Great these diseases took place during that period;						
	(1)either	○ [II.1.2.	collection o		h disease during the 12 months immediately prior to) and did not carry out vaccination against foot-and-mouth						
	(1)or	○ [II.1.2.	collection of disease dur was vaccin animal of s days prior	has not been free from foot-and-mouth disease during the 12 months immediately prior to collection of the ova(1)/embryos(1) and/or carried out vaccination against foot-and-mouth disease during that period and the donor females come from holdings on which no animal was vaccinated against foot-and-mouth disease during 30 days prior to collection and no animal of susceptible species showed clinical signs of foot-and-mouth disease during the 30 days prior to, and at least 30 days after, the ova(1)/embryos(1) were collected and the ova(1)/embryos(1) were not subjected to penetration of zona pellucida;]							
	II.2.	The ova(1)	/embryos(1)	to be exported:							
		II.2.1.	there was r		processed on premises withing -mouth disease, vesicular sto- neir collection;						
		II.2.2.	incidence o	were stored at all times on approved premises within a 10-km radius of which there was no incidence of foot-and-mouth disease, vesicular stomatitis or Rift Valley fever from the time of their collection until 30 days thereafter;							
		II.2.3.	were collected(1)/produced(1) by the team described in box I.11., which has been approved and supervised in accordance with the conditions for the approval and supervision of embryo collection teams and embryo production teams laid down in Chapter I(III)of Annex D to Directive 92/65/EEC;								
		II.2.4.	meet the co	onditions for ova and en	nbryos laid down in Chapter l	(II(II) of Annex D to Directive					
		II.2.5.	come from	the donor females of ov	rine (1)/caprine(1) species wh	ich:					
	(1)	either	○ [II.2.5.1.		gue virus-free country or zon of the ova(1)/embryos(1);]	e for at least 60 days prior to,					
	(1)	or	○ [II.2.5.1.	were kept during a blu zone;]	etongue virus seasonally free	period in a seasonally free					
	(1)	or		collection of the ova(1)	•						
	(1)	or	○ [II.2.5.1.	virus serogroup, carrie and Vaccines for Terre	al test for the detection of ant ed out in accordance with the strial Animals between 21 an and giving negative results;]	Manual of Diagnostic Tests					
	(1)	or	○ [II.2.5.1.	accordance with the M Animals on a blood sar	entification test for bluetongu anual of Diagnostic Tests and nple taken on the day of the c ing and giving negative resul	Vaccines for Terrestrial ova(1)/embryos(1) collection					
			II.2.5.2.	contact with animals of system and according to following diseases has	ledge do not come from holdi f a holding, in which, based o to the written declaration mad been clinically detected withi o collection of the ova(1)/emb	n the official notification de by the owner, any of the in the periods referred to in					
				Mycoplasm	agalactia of sheep or goats (Macapricolum, Mycoplasma mithin the last six months;						

de 2 / 5

`	THOTTHO	CUE ONION	•					(٧٥.0)		
	II. Gesundh	eitsinformatior	en							
				(b)	paratuberc months;	ulosis and caseous lym	phadenitis	s, within the last 12		
				(c)	pulmonary	adenomatosis, within	the last thi	ree years;		
_			either	 [(d) Maedi Visna for sheep or caprine viral arthritis/encephalit goats, within the last three years;] 						
ertification	(1)(4)	(1)	or	 [(d) Maedi Visna for sheep or caprine viral arthritis/encephalitis for goats, within the last 12 months, and all the infected animals were slaughtered and remaining animals subsequently reacted negativ to two tests carried out at least six months apart;] 						
7 II: C			II.2.5.3.	showed no	_	s of disease on the day	of the ova	n(1)/embryos(1)		
Pa	(1)(4)	either	○ [II.2.5.4.		originate from the region described in Box I.8., which has been recognised as officially brucellosis (B. melitensis)-free, and]					
	(1)	or	○ [II.2.5.4.	have belonged to a holding which has obtained and maintained its officially brucellosis (B. melitensis)–free status in accordance with Directive 91/68/EEC, and]						
	(1)	or	○ [II.2.5.4.	originate from a holding, where in respect of brucellosis (B. melitensis) all susceptible animals have been free from any clinical or any signs of this disease for the last 12 months, none of the ovine and caprine animals have been vaccinated against this disease, save those vaccinated with Rev. 1 vaccine more than two years ago, and all ovine and caprine animals over six months of age have been subjected to at least two tests (3), carried out with negative results on samples taken on (date) and on (date) at least six months apart, the latter being within 30 days prior to collection of the ova(1)/embryos(1),]						
	and		have not be		•	holding of lower status	s;			
	(1)	either	○ [II.2.5.5.	have remained in the exporting country for at least the past six months prior collection of the ova(1)/embryos(1) to be exported;]				past six months prior to		
	(1)	or	○ [II.2.5.5.	complied ova/embry been impo	with the anin yos(1)which a	exporting country at le	pplying to to the Gre			
			II.2.5.6.		kept continu s are fulfilled:	ously since birth in a c	country wh	nere the following		
				II.2.5.6.1.	classical sci	apie is compulsorily n	otifiable;			
				II.2.5.6.2.	an awarene	ess, surveillance and m	onitoring :	system is in place;		
				II.2.5.6.3.		aprine animals affecte tely destroyed;	d with clas	ssical scrapie are killed		
				II.2.5.6.4.	greaves of r	ruminant origin has be	en banned	meat-and-bone meal or l and effectively of at least the last seven		
	(1)	either	○ [II.2.5.7.	. have been kept continuously for the last three years before the collection of the embryos to be exported in a holding or holdings which has/have been complyir for the last three years before the collection of the embryos to be exported with the requirements laid down in points (a) to (f) of point 1.3. of Section A of Chapter A of Annex 8 to Regulation (EC) No 999/2001;]				has/have been complying ryos to be exported with		
	(1)	or	○ [II.2.5.7.	are ovine	animals and	the embryos				
			(1)	either		e ARR/ARR prion prote		-		
			(1)	or	o [carry at January 201		nd were co	llected after the date of 1		
	[II.2.6. were collected(1)/produced(1)in the exporting country,									

EL	JROPÄISCI	1E UNION						(v3.0)		
	II. Gesundhei	tsinformatione	n							
	(1)	either	o [II.2.6.1.	which acc (EHD);]	cording to off	icial findings is free fro	om epiz	zootic haemorrhagic disease		
uo	(1)(5)	or	 [II.2.6.1. in which according to official findings the following serotypes of epizootic haemorrhagic disease (EHD) exist: and the donor females of ovine(1)/caprine(1) species were subjected with negative results in each case the following tests carried out in an approved laboratory: 							
Part II: Certification			(1)	either	serogroup, not more tl	carried out on sample nan 12 months apart p	s of blo rior to	of antibody to the EHD virus ood taken on two occasions and not less than 21 days t of ova(1)/embryos(1);]		
Part II:			(1)	or	serogroup, more than	carried out on sample 60 days throughout th s after the final collect	es of blo e collec	of antibody to the EHD virus ood taken at intervals of not ction period and between 21 this consignment of		
			(1)	or	collected at days, if car carried out	commencement and cried out as virus isolat	conclus ion test reaction	d out on samples of blood sion of, and at least every 7 t, or at least every 28 days, if n, during collection for this		
		II.2.7.		_	d(1)/produced(1) after the date on which the embryo collection team was the competent authority of the exporting country;					
		II.2.8.	were processed and stored under approved conditions for at least 30 days immediately after their collection(1)/production(1) and transported under conditions for ova and embryos laid down in Chapter III(II) of Annex D to Directive 92/65/EEC							
		II.2.9.		ance with the requirements (II) of Annex D to Directive						
	(1) 🗆	[II.2.10.	the consignment consists of embryos of the ovine or caprine species which were conceived by artificial insemination (1)/as a result of in vitro fertilisation (1) using semen coming from semen collection centres approved (7) in accordance with:							
	(1)	either	o [II.2.10.1.			ve 92/65/EEC and locate irements of Directive		reat Britain; and the semen EC.]		
	(1)(8)or	° [II.2.10.1.	Article 17(3)(b) of Directive 92/65/EEC and located in a third country or part thereof as set out in a document relating to 'ovine and caprine semen' published on gov.uk, in accordance with Commission Decision 2010/472/EU, and the semen complies with the requirements set out in the relevant export health certificate, in the form published by the Secretary of State and amended from time to time.]							
	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).									
			•	•	•	nel Islands and Isle of	Man.			
	Part I:									
	Box reference I.6:	Person restransit con	esponsible for the load in Great Britain: this box is to be filled in only if it is a certificate for ommodity.							
	Box reference I.11:	which the	lace of origin shall correspond to the approved embryo collection team or embryo production team by which the ova/embryos were collected/produced, processed and stored; and listed in accordance with rticle 17(3)(b) of Directive 92/65/EEC.							
	Box reference I.22:	Number of packages shall correspond to the number of containers.								

Edward Chief																	
	II. Gesundheitsinformationen																
	Box reference I.23:	reference															
Part II: Certification	Box reference I.26:	Fill in according to whether it is a transit or an import certificate. nce															
	Box reference I.27:	Fill in according to whether it is a transit or an import certificate.															
Part II: C	Box reference I.28:	Species: Select amongst 'Ovis aries' or 'Capra hircus' as appropriate															
_	Category: specify if in vivo derived embryos, in vivo derived ova, in vitro produced embryos or micromanipulated embryos.																
	Donor ider	ntity shall correspond to the official identification	on of the animal.														
	Date of col	lection shall be indicated for in vivo derived em	abryos and in the following fo	ermat: dd.mm.yyyy.													
	Date of fre	ezing shall be indicated in the following format	: dd.mm.yyyy.														
	team by w	number of the team: shall correspond to the app hich the ova/embryos were collected/produced, 3)(b) of Directive 92/65/EEC.															
	Part II:																
	(1)	Delete as appropriate.	copriate.														
	(2)	Only third countries or parts thereof as set out published on gov.uk, in accordance with Comr	•	-													
	(3)	Tests shall be carried out in accordance with A	annex C to Directive 91/68/EE0	2.													
	(4)	Only for the territory appearing with the entry 'V' in column 6 of a document relating to 'live ungulates published on gov.uk, in accordance with Commission Regulation (EU) No 206/2010.(8)															
	(5)	See remarks for exporting country or part thereof as set out in a document relating to 'ovine and caprine ova and embryos' published on gov.uk, in accordance with Decision 2010/472/EU.(8)															
	(6)	Standards for EHD virus diagnostic tests are described in Chapter 2.1.3. of the WOAH (formerly OIE) Manual of Diagnostic Tests and Vaccines for Terrestrial Animals.															
	(7)	Only approved semen collection centres listed Directive 92/65/EEC.	in accordance with Article 11	l(4) and Article 17(3)(b) of													
	e and caprine ova and ne consent of the Scottish and																
EU and EFTA states approved to export animals and animal products to Great Britain - data.gov.uk The signature and the stamp must be in a different colour to that of the printing. Certifying Officer Name (in capital letters) Qualification and title																	
										Datum der Unterzeichnung Unterschrift Stempel							

de **5** / 5