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►<u>B</u>

COMMISSION DECISION

of 26 August 2010

on imports of semen, ova and embryos of animals of the ovine and caprine species into the Union

(notified under document C(2010) 5780)

(Text with EEA relevance)

(2010/472/EU)

(OJ L 228, 31.8.2010, p. 74)

Amended by:

Official Journal

		No	page	date
► <u>M1</u>	Commission Implementing Decision 2012/411/EU of 17 July 2012	L 192	16	20.7.2012
► <u>M2</u>	Commission Regulation (EU) No 519/2013 of 21 February 2013	L 158	74	10.6.2013
► <u>M3</u>	Commission Implementing Decision 2013/470/EU of 20 September 2013	L 252	32	24.9.2013
► <u>M4</u>	Commission Implementing Decision 2014/802/EU of 14 November 2014	L 331	28	18.11.2014
► <u>M5</u>	Commission Implementing Decision (EU) 2016/2002 of 8 November 2016	L 308	29	16.11.2016

Corrected by:

▶<u>C1</u> Corrigendum, OJ L 331, 1.12.2012, p. 57 (2012/411/EU)

COMMISSION DECISION

of 26 August 2010

on imports of semen, ova and embryos of animals of the ovine and caprine species into the Union

(notified under document C(2010) 5780)

(Text with EEA relevance)

(2010/472/EU)

Article 1

Subject matter

This Decision sets out a list of third countries or parts thereof from which Members States are to authorise the importation into the Union of consignments of semen, ova and embryos of animals of the ovine and caprine species.

It also lays down certification requirements for the importation of those commodities into the Union.

Article 2

Imports of semen

Member States shall authorise imports of consignments of semen of animals of the ovine and caprine species provided that they comply with the following conditions:

- (a) they come from a third country or part thereof listed in Annex I;
- (b) they come from an approved semen collection or storage centre listed in accordance with Article 17(3)(b) of Directive 92/65/EEC;
- (c) they are accompanied by a health certificate drawn up in accordance with the following model health certificates set out in Part 2 of Annex II, and completed in accordance with the explanatory notes set out in Part 1 of that Annex:
 - (i) model 1 as set out in Section A, for consignments of semen dispatched from an approved semen collection centre of origin of the semen;
 - (ii) model 2 as set out in Section B, for consignments of semen dispatched from an approved semen storage centre.

However, where specific certification requirements are laid down in bilateral agreements between the Union and third countries, those requirements shall apply.

(d) they comply with the requirements set out in the health certificates referred to in point (c).

Article 3

Imports of ova and embryos

Member States shall authorise imports of consignments of ova and embryos of animals of the ovine and caprine species provided that they comply with the following conditions:

(a) they come from a third country or part thereof listed in Annex III;

- (b) they come from an approved embryo collection or production team listed in accordance with Article 17(3)(b) of Directive 92/65/EEC;
- (c) they are accompanied by a health certificate drawn up in accordance with the model set out in Part 2 of Annex IV, and completed in accordance with the explanatory notes set out in Part 1 of that Annex.

However, where specific certification requirements are laid down in bilateral agreements between the Union and third countries, those requirements must apply.

(d) they comply with the requirements set out in the health certificate referred to in point (c).

Article 4

General conditions concerning the transport of consignments of semen, ova and embryos to the Union

1. Consignments of semen, ova and embryos of animals of the ovine and caprine species shall not be transported in the same container as other consignments of semen, ova and embryos that:

(a) are not intended for introduction into the Union, or

(b) are of a lower health status.

2. During transport to the European Union, consignments of semen, ova and embryos shall be placed in closed and sealed containers and the seal must not be broken during the transport.

Article 5

Repeal

Decision 2008/635/EC is repealed.

Article 6

Transitional provisions

For a transitional period until 31 August 2011, Member States shall authorise imports from third countries of stocks of the following commodities:

- (a) semen of animals of the ovine and caprine species which were collected, processed and stored in accordance with Directive 92/65/EEC by 31 August 2010 and which are accompanied by a health certificate issued not later than 31 May 2011 in accordance with the model set out in Annex II to Decision 2008/635/EC.
- (b) ova and embryos of animals of the ovine and caprine species which were collected or produced, processed and stored in accordance with Directive 92/65/EEC by 31 August 2010 and which are accompanied by a health certificate issued not later than 31 May 2011 in accordance with the model set out in Annex VI to Decision 2008/635/EC.

Article 7

Applicability

Article 8

Addressees

This Decision is addressed to the Member States.

ANNEX I

List of third countries or parts thereof from which Member States are to authorise imports of consignments of
semen of animals of the ovine and caprine species

			Remarks		
	ISO Code	Name of the third country	Description of the territory (<i>if appropriate</i>)	Additional guarantees	
	AU	Australia		The additional guarantee as regards testing set out in point II.4.9.1 of the model health certificate set out in Section A of Part 2 of Annex II is compulsory.	
	СА	Canada		The additional guarantee as regards testing set out in point II.4.9.1 of the model health certificate set out in Section A of Part 2 of Annex II is compulsory.	
	СН	Switzerland (1)			
	CL	Chile			
	GL	Greenland			
▼ <u>M2</u>					
▼ <u>M1</u>					
	IS	Iceland			
	NZ	New Zealand			
	PM	Saint Pierre and Miquelon			
	US	United States		The additional guarantee as regards testing set out in point II.4.9.1 of the model health certificate set out in Section A of Part 2 of Annex II is compulsory.	

(1) Certificates in accordance with the Agreement between the European Community and the Swiss Confederation on Trade in Agricultural Products as approved by Decision 2002/309/EC, Euratom of the Council, and of the Commission as regards the Agreement on Scientific and Technological Cooperation of 4 April 2002 on the conclusion of seven Agreements with the Swiss Federation (OJ L 114, 30.4.2002, p. 1).

▼<u>M1</u>

ANNEX II

PART 1

Explanatory notes for the certification

(a) The health certificates shall be issued by the competent authority of the exporting third country, in accordance with the model set out in Part 2 of Annex II.

If the Member State of destination requires additional certification requirements, attestations to certify that those requirements are fulfilled shall be also incorporated in the original form of the health certificate.

- (b) The original of the health 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) Where the model health 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 certifying officer, or completely deleted from certificate.
- (d) The health 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 European Union and of the Member State of destination. However, those Member States may authorise the certificate to be drawn up in the official language of another Member State, and accompanied, if necessary, by an official translation.
- (e) If for the reasons of identification of the items of the consignment (schedule in Box I.28 of the model health certificate), additional sheets of paper are attached to the health certificate, those sheets of paper shall also be considered as forming part of the original of the health certificate by application of the signature and stamp of the certifying officer, on each of the pages.

- (f) When the health certificate, including additional schedules referred to in (e), 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 on the top of the pages.
- (g) The original of the health certificate must be completed and signed by an official veterinarian the last working day prior to loading of the consignment for exportation to the European Union. The competent authorities of the exporting third country shall ensure that certification requirements equivalent to those laid down in Council Directive 96/93/EC (¹) are followed.

The colour of the signature and the stamp of the official veterinarian shall be different to that of the printing on the health certificate. This requirement also applies to stamps other than those embossed or watermarks.

- (h) The original of the health certificate must accompany the consignment until it reaches the border inspection post of introduction into the European Union.
- (i) The certificate reference number referred to in Box I.2 and Box II.a of the model health certificate must be issued by the competent authority of the exporting third country.

PART 2

Model health certificates for imports of consignments of semen of animal of the ovine and caprine species

▼<u>M5</u>

Section A

MODEL 1 — Health certificate for semen dispatched from an approved semen collection centre of origin of the semen

COUNTRY:

Veterinary certificate to EU

	l.1.	Consignor				I.2. Certificate reference No I.2.a.						
		Name Address					I.3. Central competent authority					
		Address					I.4. Local competent authority					
		Tel.	əl.									
	1.5.	Consignee	Consignee					Person respo	nsible for th	ne load	d in EU	
t		Name	Jame					Name				
nmen		Address	Address					Address				
onsig		Postal code	Postal code					Postal code				
ned c		Tel.						Tel.				
Part I: Details of dispatched consignment	1.7.	Country of origin	ISO code	l.8.	Region of origin	Code	I.9.	Country of destination	ISO code	l.10.	Region of destination	Code
Details	111	Place of or	iain				112	Place of desti	nation			
art I: I	1. 1 1.		igiri				1.12.		nation			
ã		Name		Ap	proval nun	nber		Name				
		Address					Address					
		Name		Ap	proval nun	nber						
		Address						Postal code				
		Name		Ap	proval nun	nber						
		Address										
	l.13.	Place of loa	ading				I.14. Date of departure					
	l.15.	Means of t	ransport				l.16.	Entry BIP in E	U			
		Aeroplane		Ship E] Rai	lway						
		wagon 🗖	. —		_		I.17.					
		Road vehic		Othe	er 🗖				_			
		Documenta		nces								
	1 1 0	Description							110 000		ty code (HS co	do)
	1.10.	Description		louity					1.19. Con		5 11 99 85	ue)
								L		1.2	0. Quantity	

I.21.	I.22. Number of packages
I.23. Seal/Container No	1.24.
I.25. Commodities certified for: Artificial reproduction	
I.26. For transit through EU to third country I I.27. For import or admission Third country ISO code ISO code	n into EU 🛛
I.28. Identification of the commodities	
(Scientific name) collection number	roval Quantity er of the ntre

_	COUNTRY			Ovine and ca	aprine semen — Section /				
	II.	Health in	formation	II.a. Certificate reference No	II.b.				
_	I, the undersigned, official veterinarian, hereby certify that:								
	II.1.	The expo		ne of exporting country) (²)					
		II.1.1.	contagious caprine immediately prior to c	rinderpest, peste des petits rumina pleuropneumonia and Rift Valley fev collection of the semen to be exported a accination against these diseases took p	ver during the 12 month and until its date of dispate				
 contagious caprine pleuropneumonia and Rift Valley fever during the immediately prior to collection of the semen to be exported and until its date to the Union and no vaccination against these diseases took place during that II.1.2. has been free from foot-and-mouth disease during the 12 months immediately collection of the semen to be exported and until its date of dispatch to the Ur vaccination against this disease took place during that period. 									
	II.2.		en collection centre de and stored:	escribed in Box I.11 and at which the	semen to be exported wa				
		II.2.1.		s for the approval of semen collec ex D to Directive 92/65/EEC;	tion centres laid down i				
		II.2.2.		pervised in accordance with the cond nd storage centres laid down in Chaj					
	II.3.	The ovine	e (¹)/caprine (¹) animals	standing at the semen collection centre	:				
		II.3.1.	prior to their stay in th	e quarantine accommodation described	in point II.3.3,				
	(¹) (⁴) either	[II.3.1.1.		originate from the territory described in Box I.8, which has been recognised as officially brucellosis (<i>B. melitensis</i>)-free,]					
	(¹) or	[II.3.1.1.	have belonged to a holding which has obtained and maintained its officially brucellosis (<i>B. melitensis</i>)-free status in accordance with Directive 91/68/EEC,]						
	(¹) or	[II.3.1.1.	originate from a holding, where in respect of brucellosis (<i>B. melitensis</i>) all susceptible animals have been free from 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 our with negative results on samples taken on (<i>date</i>) and on						
	and		have not been kept pr	reviously in a holding of a lower status;					
		II.3.1.2.		tinuously for at least 60 days on a is (<i>Brucella ovis</i>) has been diagnosed ir					
	(¹) and		[they are animals of the ovine species and have undergone during the 60 days prior their stay in the quarantine accommodation described in point II.3.3 a compleme fixation test, or any other test with an equivalent documented sensitivity and specificity, detect contagious epididymitis with result of less than 50 ICFTU/ml;]						
II.3.1.3. to the best of my knowledge do not come from holdings and have not b with animals of a holding, in which, based on the official notification according to the written declaration made by the owner, any of the follow has been clinically detected within the periods referred to in points (a) to (or stay in the quarantine accommodation described in point II.3.3.									
				ilactia of sheep or goats (Mycoplasm coplasma mycoides var. mycoides "lar					
			(b) paratuberculosis	s and caseous lymphadenitis, within the	last 12 months,				
			(c) pulmonary aden	nomatosis, within the last three years;					

[5			

COUNTRY			Ovine and ca	prine semen — Section A
II.	Health in	formation	II.a. Certificate reference No	II.b.
	(¹) either	[(d) Maedi/Visna for s three years;]	sheep or caprine viral arthritis/encephali	itis for goats, within the last
	(¹) or	12 months, and a	sheep or caprine viral arthritis/encephali all the infected animals were slaughter icted negatively to two tests carried out	red and remaining animals
	II.3.2.		ollowing tests carried out on a blood s commencement of the period of quaran	
		— brucellosis (<i>B. m</i> Annex C to Direc	<i>elitensis</i>), with negative results in eac tive 91/68/EEC;	h case in accordance with
		results in each c	dymitis (<i>Brucella. ovis</i>), in the case of ase in accordance with Annex D to D equivalent documented sensitivity and	irective 91/68/EEC, or any
		 border disease i Directive 92/65/E 	in accordance with point 1.4(c) of Ch EC;	apter II(II) of Annex D to
	II.3.3.		arantine isolation period of at least ically approved for the purpose by the	
	II.3.3.1.	only animals of at le accommodation;	east the same health status were $\mathfrak p$	present in the quarantine
	II.3.3.2.	by the competent auth	argone the following tests, carried out the original of the exporting country on same als were admitted to the quarantine accounts of the second of the seco	ples taken not earlier than
		— brucellosis (<i>B. m.</i> Annex C to Direc	nelitensis) with negative results in eacl tive 91/68/EEC;	h case in accordance with
		results in each c	dymitis (<i>Brucella ovis</i>), in the case of ase in accordance with Annex D to D equivalent documented sensitivity and	irective 91/68/EEC, or any
		 border disease in 92/65/EEC; 	accordance with point 1.6 of Chapter I	I(II) of Annex D to Directive
	II.3.4.	have undergone at leas	st once a year the routine tests for:	
		— brucellosis (<i>B. m.</i> Annex C to Direc	nelitensis) with negative results in eacl tive 91/68/EEC;	h case in accordance with
		results in each c	dymitis (<i>Brucella ovis</i>), in the case of ase in accordance with Annex D to D equivalent documented sensitivity and	irective 91/68/EEC, or any
		— border disease Directive 92/65/E	in accordance with point 5(c) of Cha EC.	apter II(II) of Annex D to
11.4.	The seme	en to be exported was ob	otained from donor rams (1)/bucks (1) wh	nich:
	II.4.1.	were admitted to the a the centre veterinarian.	approved semen collection centre with	the express permission of
	II.4.2.		s of disease on the day of admission n the day the semen was collected;	n to the approved semen
(¹) either	[11.4.3.	have not been vaccina collection of the semen	ted against foot-and-mouth disease du ;]	ring the 12 months prior to

II.	Health in	formation	II.a. Certificate reference No	II.b.			
(¹) or	[11.4.3.	have been vaccinated against foot-and-mouth disease at least 30 days prior to the collection, and 5 % (with a minimum of five straws) of each collection have been submitted to a virus isolation test for foot-and-mouth disease with negative results;]					
	II.4.4.		approved semen collection centre for ely prior to collection of the semen, in	•			
	II.4.5.		lly after their entry to the quarantine ac d including the day of semen collection				
	II.4.6.	have been kept at appro	oved semen collection centres:				
	II.4.6.1.	collection of the semen the date of dispatch, a	from foot-and-mouth disease for at le and 30 days after collection or, in the nd which are situated in the centre of as been no case of foot-and-mouth dis semen;	case of fresh semen, unt an area of 10 kilometre			
	II.4.6.2.	ending 30 days after co	during the period commencing 30 da llection of the semen or, in the case of ellosis (<i>B. melitensis</i>), contagious ep	fresh semen, until the dat			
(¹) either	[11.4.7.	have remained in the exponent	xporting country for at least the past siv orted;]	< months prior to collectio			
(¹) or	[11.4.7.	during the last six months prior to collection of the semen they complied with the animal health conditions applying to donors of the semen which is intended for export to the Union and they have been imported into the exporting country at least 30 days prior to collection of the semen from					
(¹) either	[11.4.8.	were kept in a blueton during, collection of the	gue virus-free country or zone for at le semen;]	east 60 days prior to, an			
(¹) or	[11.4.8.		etongue virus seasonally free period in , and during collection of the semen;]	a seasonally free zone fo			
(¹) or	[11.4.8.	were kept in a vector-p collection of the semen;	protected establishment for at least 60]	days prior to, and durin			
(¹) or	[11.4.8.	were subjected to a serological test for the detection of antibody to the bluetongue virus group, carried out in accordance with the OIE Manual of Diagnostic Tests and Vaccines for Terrestrial Animals, with negative results, on blood samples taken at least every 60 days throughout the collection period and between 21 and 60 days after the fina collection for this consignment of semen;]					
(¹) or	[11.4.8.	were subjected to an agent identification test for bluetongue virus, carried out in accordance with the Manual of Diagnostic Tests and Vaccines for Terrestrial Animals with negative results on blood samples taken at commencement and final collection for this consignment of semen and at least every 7 days (virus isolation test) or at least every 28 days (PCR test) during collection for this consignment of semen;]					
(¹) (⁵) either	[11.4.9.	were resident in the exercise epizootic haemorrhagic	<pre>kporting country which according to of disease (EHD);]</pre>	ficial findings is free fror			
(¹) or	[11.4.9.	serotypes of epizootic h	porting country in which according to of aemorrhagic disease (EHD) exist: gative results in each case to:				
	(¹) either	approved laboratory o	or the detection of antibody to the EHE n samples of blood taken on two of to and not less than 21 days after t	occasions not more tha			

COUNTRY				Ovir	ne and cap	orine semen — Section A		
II.	Health in	formation	II.a.	Certificate reference	e No	II.b.		
	(¹) or	[a serological test (⁶) for the detection of antibody to the EHDV group, carried out in an approved laboratory on samples of blood taken at intervals of not more than 60 days throughout the collection period and between 21 and 60 days after the final collection for this consignment of semen.]]						
	(¹) or	taken at commenceme	nt and	l conclusion of, and a	at least eve	ratory on samples of blood ery 7 days (virus isolation n for this consignment of		
	II.4.10.	have been kept continu fulfilled:	ously	since birth in a countr	ry where th	ne following conditions are		
	II.4.10.1.	classical scrapie is com	classical scrapie is compulsorily notifiable;					
	II.4.10.2.	an awareness, surveilla	nce ai	nd monitoring system is	s in place;			
	II.4.10.3.	ovine and caprine anin destroyed;	vine and caprine animals affected with classical scrapie are killed and completely estroyed;					
	II.4.10.4.	the feeding to ovine and caprine animals of meat-and-bone meal, or greaves of ruminant origin, as defined in the OIE Terrestrial Animal Health Code, has been banned and effectively enforced in the whole country for a period of at least the last seven years;						
(¹) either	[11.4.11.	collection of the semen during that period all t Chapter A of Annex VIII	to be he ree to Re men e	exported in a holding quirements set out in gulation (EC) No 999/ collection centre that	or holding points 1.3 2001, exce complied d	s preceding the date of the gs which has/have fulfilled (a) to (f) of Section A of ppt during the period when luring that period with the ection;]		
(¹) or	[11.4.11.	are ovine animals of AR	R/AR	R prion protein genoty	pe.]			
II.5.	The seme	en to be exported:						
	II.5.1.	was collected after the competent authority of t			ollection ce	entre was approved by the		
	II.5.2.					d in accordance with the) of Annex D to Directive		
	II.5.3.		to be	e subject to trade laid	down in po	in accordance with the pint 1.4 of Chapter III(I) of ated in Box I.23.		
(¹) either	[II.6.	No antibiotics were add	ed to t	he semen.]				
(¹) or	[II.6.	The following antibioti concentration in the fina				as added to produce a		
]			
Notes								
Part I:								
Box I.6:	Person res commodity		<i>EU</i> : tl	nis box is to be filled	in only if if	t is a certificate for transit		
Box I.11:	collected a		with A	rticle 17(3)(b) of Dire	ctive 92/65	in which the semen was /EEC on the Commission tm		
Box I.22:	Number of	packages shall correspor	nd to t	he number of containe	rs.			

COUNTRY		Ovine and	d caprine semen — Section A
II.	Health information	II.a. Certificate reference No	II.b.
Box I.23:	Identification of container and se	eal number shall be indicated.	
Box I.26:	Fill in according to whether it is	a transit or an import certificate.	
Box I.27:	Fill in according to whether it is	a transit or an import certificate.	
Box I.28:	Species: select amongst "Ovis a	aries" or "Capra hircus" as appropriate.	
	Donor identity shall correspond	to the official identification of the anima	al.
	Date of collection shall be indica	ated in the following format: dd.mm.yyy	у.
	Approval number of the centre centre indicated in Box I.11.	e shall correspond to the approval nu	umber of the semen collection
Part II:			
(1) Delet	te as necessary.		
(²) Only	third countries listed in Annex I to	Decision 2010/472/EU.	
(³) Tests	s shall be carried out in accordanc	e with Annex C to Directive 91/68/EEC	;.
	for the territory appearing with the No 206/2010 (OJ L 73, 20.3.2010	e entry "V" in column 6 of Part 1 of Anr , p. 1.).	nex I to Commission Regulation
(⁵) See	remarks for exporting country con	cerned in Annex I to Decision 2010/472	2/EU.
	dards for EHD virus diagnostic te s and Vaccines for Terrestrial Anir	ests are described in Chapter 2.1.7 of nals.	the OIE Manual of Diagnostic
(⁷) Inser	t names and concentrations.		
— The s	signature and the stamp must be i	n a different colour to that of the printin	g.
Official ve	terinarian		
	Name (in capital letters):	Qu	alification and title:
	Date:	Sig	nature:
	Stamp:		

COUNTRY

Section B

MODEL 2 - Health certificate for semen dispatched from an approved semen storage centre

cou	NTRY	′:				Veterinary cert	ificate to EU	
	l.1.	Consignor	1.2.	Certificate reference I	No	1.2.a.		
		Name						
		Address	1.3.	Central competent au	thority			
		Tel.	1.4.	Local competent auth	ority			
	1.5.	I.5. Consignee I.6		I.6 Person responsible for the load in EU				
jt		Name		Name				
l E		Address		Address				
sig								
cor	Postal code Tel.			Postal code Tel.				
ber		161.		161.				
dispached consignment		100			100			
disl	1.7.	Country of origin ISO code I.8. Region of origin Code	1.9.	Country of destination	ISO I. code	10. Region of destination	Code	
ď				destination		destination	1	
ails	1 4 4	Diago of evicin						
Part I: Details of	1. 1 1.	Place of origin	1.12.	Place of destination				
μ		Name Approval number		Name				
Pai		Address		Address				
		Name Approval number						
		Address		Postal code				
		Name Approval number						
		Address						
	1.13.	Place of loading	1.14.	Date of departure				
	I.15.	Means of transport	I.16.	Entry BIP in EU				
		Aeroplane 🗌 Ship 🗌 Railway wagon 🗌						
		Road vehicle Other O	I.17. No(s) of related original certificates					
		Identification		····(-) -· ·······				
		Documentary references						
	l.18.	Description of commodity		I.19. Comm	odity code	(HS code)		
					05 1	1 99 85		
					1.20. Qu	lantity		
	1.21.				1.22. Nu	mber of packages		
	1.23.	Seal/container No	1.24.					
	125	Commodities certified for:						
		_						
		Artificial reproduction						
	1.26.	For transit through the EU to a third country	1.27.	For import or admission	on into the	EŲ		
		Third country ISO code						
	1.28.	Identification of the commodities						
			of	llection A	pproval and	mber	Quantity	
		Species Breed Donor identity Date (scientific name)		A A A	pproval nur of the cen		Quantity	

COUNTRY	Υ:					
II. I	Health i	information		II.a. Certificate reference No	II.b.	
I, the und	dersigne	ed official ve	eterinarian of	e of exporting country) (²)		hereby certify the
		II.1.	The centre (3) described in Box I.11 at v	which the semen to be exported to	the Europe	an Union was stored:
(¹) either		[1].1.1.	meets the conditions laid down in Chapt	er I(I)(1) of Annex D to Directive S	2/65/EEC;	
	and	II.1.2.	is operated and supervised in accordance 92/65/EEC.]	ce with the conditions laid down ir	n Chapter I(I	I)(1) of Annex D to Direct
(1) or		[.1.1.	meets the conditions laid down in Chapt	er I(I)(2) of Annex D to Directive S	2/65/EEC;	
	and	II.1.2.	is operated and supervised in accordance 92/65/EEC.]	ce with the conditions laid down ir	1 Chapter I(I	I)(2) of Annex D to Direct
		II.2.	The semen to be exported to the Europe	ean Union:		
		II.2.1.	has been collected, processed and storr approved semen collection centre (⁴) ope of Annex D to Directive 92/65/EEC, and	erated and supervised in accordance		
		(1) either	[located in the exporting country;]			
		(¹) and/or	[located in			
		and	has been imported to the exporting cour caprine species into the European Unior			ports of semen of ovine a
		II.2.2.	was moved to the centre described in Pa to Decision 2010/472/EU (⁶);]	rt I.11 under conditions at least as	strict as in S	ection A of Part 2 of Anne
		II.2.3.	was stored under conditions which satisf	fy the terms of Annex D to Directi	/e 92/65/EE	С;
		II.2.4.	was sent to the place of loading in a se Directive 92/65/EEC and bearing the nur		point 1.4 of	i Chapter III(I) of Annex D
Notes						
Part I:						
Box I.11:	place	of origin sh	all correspond to the approved semen sto	rage centre of dispatch of the sem	ien.	
Box I.17:	above	from the ap	o the serial number of the individual official proved semen collection centre of its origin or the officially endorsed copies of there	to the centre described in Box I.1	1. The origin	
Box 1.22:	numbe	er of packag	es shall correspond to the number of con	tainers.		
Box 1.23:	identif	ication of co	ntainer and seal number shall be indicate	d.		
Box 1.28:	donor	identity sha	Il correspond to the official identification o	f the animal.		
	date o	of collection	shall be indicated in the following format:	dd/mm/yyyy.		
	appro	val number o	of the centre shall correspond to the approv	val number of the approved semen	collection ce	antro in which the comon w

II.	Health information	II.a. Certificate reference No	II.b.
Part	II:		
(¹) C	Delete as necessary.		
(²) C	Only third countries listed in Annex I to Decision 2010/472/EU.		
	Only approved semen collection or storage centres listed in acc vebsite:	ordance with Article 17(3)(b) of Dire	ctive 92/65/EEC on the Commissior
h	http://ec.europa.eu/food/animal/semen_ova/ovine/index_en.htm		
	Only approved semen collection centres listed in accordance wi vebsites:	th Article 11(4) and 17(3)(b) of Dire	ctive 92/65/EEC on the Commission
	http://ec.europa.eu/food/animal/approved_establishments/establishm http://ec.europa.eu/food/animal/semen_ova/ovine/index_en.htm	nents_vet_field_en.htm	
(⁵) C	Only third countries listed in Annex I to Decision 2010/472/EU and	d the EU Member States.	
) a	The original(s) of the document(s) or the health certificate(s) or the above from the approved semen collection centre in which the so dispatch described in Box I.11 must be attached to this certificate.	emen was collected to the approved	
Offici	ial veterinarian (*)		
Ν	Name (in capital letters):		Qualification and title:
C	Date:		Signature:
S	Stamp:		

ANNEX III

				Remarks
	ISO Code	Name of the third country	Description of the territory <i>(if appropriate)</i>	Additional guarantees
	AU	Australia		The additional guarantee as regards testing set out in point II.2.6.1 of the model health certificate set out in Part 2 of Annex IV is compulsory.
	СА	Canada		The additional guarantee as regards testing set out in point II.2.6.1 of the model health certificate set out in Part 2 of Annex IV is compulsory.
	СН	Switzerland (1)		
	CL	Chile		
	GL	Greenland		
▼ <u>M2</u>				
▼ <u>M1</u>				
	IS	Iceland		
	NZ	New Zealand		
	РМ	Saint Pierre and Miquelon		
	US	United States		The additional guarantee as regards testing set out in point II.2.6.1 of the model health certificate set out in Part 2 of Annex IV is compulsory.

List of third countries or parts thereof from which Member States are to authorise imports of consignments of ova and embryos of animals of the ovine and caprine species

(¹) Certificates in accordance with the Agreement between the European Community and the Swiss Confederation on Trade in Agricultural Products as approved by Decision 2002/309/EC.

▼<u>M1</u>

ANNEX IV

PART 1

Explanatory notes for the certification

(a) The health certificates shall be issued by the competent authority of the exporting third country, in accordance with the model set out in Part 2 of Annex IV.

If the Member State of destination requires additional certification requirements, attestations to certify that those requirements are fulfilled shall be also incorporated in the original form of the health certificate.

- (b) The original of the health 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) Where the model health 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 certifying officer, or completely deleted from certificate.
- (d) The health 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 European Union and of the Member State of destination. However, those Member States may authorise the certificate to be drawn up in the official language of another Member State, and accompanied, if necessary, by an official translation.
- (e) If for the reasons of identification of the items of the consignment (schedule in Box I.28 of the model health certificate), additional sheets of paper are attached to the health certificate, those sheets of paper shall also be considered as forming part of the original of the health certificate by application of the signature and stamp of the certifying officer, on each of the pages.

- (f) When the health certificate, including additional schedules referred to in (e), 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 on the top of the pages.
- (g) The original of the health certificate must be completed and signed by an official veterinarian the last working day prior to loading of the consignment for exportation to the European Union. The competent authorities of the exporting third country shall ensure that certification requirements equivalent to those laid down in Council Directive 96/93/EC (¹) are followed.

The colour of the signature and the stamp of the official veterinarian shall be different to that of the printing on the health certificate. This requirement also applies to stamps other than those embossed or watermarks.

- (h) The original of the health certificate must accompany the consignment until it reaches the border inspection post of introduction into the European Union.
- (i) The certificate reference number referred to in Box I.2 and Box II.a of the model health certificate must be issued by the competent authority of the exporting third country.

^{▼&}lt;u>B</u>

^{(&}lt;sup>1</sup>) OJ L 13, 16.1.1997, p. 28.

▼<u>M4</u>

PART 2

Model health certificate for imports of consignments of ova and embryos of animals of the ovine and caprine species

						erinary certific	suic io
1.1.	Consignor	1.2.	Certificate refere	nce No	1.2	.a.	
ĺ	Name						
	Address	1.3.	Central compete	nt authority			
	Tel.	I.4. Local competent authority					
1.5.	Consignee	1.6.	Person responsil	ble for the load	in EU		
1	Name		Name				
	Address		Address				
l l	Postal code		Postal code				
l l	Tel.		Tel.				
1.7.	Country of ISO code I.8. Region of Code origin origin	1.9.	Country of destination	ISO code	I.10.	Region of destination	Cod
1.11.	Place of origin	1.12.	Place of destinat	ion			_
1	Name Approval number		Name				
l l	Address		Address				
1	Name Approval number						
1	Address		Postal code				
ĺ	Name Approval number						
ĺ	Address						
l.13.	Place of loading		Date of departure	Э			
145	Manage	1.10	Esta DID is Ell				
I.15.	Means of transport	I.16.	Entry BIP in EU				
ĺ	Aeroplane 🛛 Ship 🖾 Railway wagon 🗖						
ĺ							
1	Road vehicle U Other U	I.17.					
ĺ							
	Documentary references						
I.18.	Description of commodity					ode (HS code)
				05 11	99 85		
					1.20.	Quantity	
1.21.					1.22.	Number of	
						packages	
1.23.	Seal/Container No				1.24.		
1.25.	Commodities certified for:						
	Artificial reproduction						
1.26.	For transit through EU to third country	I	.27. For import of	or admission ir	ito EU		
	Third country ISO code						
	ning country 100 tode						
1.28.	Identification of the commodities						
	Species Category Donor identity Date of collect	tion F	ate of freezing Ap	nroval numbe	r of the	team Ous	intity
(0.	cientific name)		ALC OF ILCOZING AL	provar numbe	1 01 1110	Qua	muty

		Health infor	mation	II.a.	Certificate refe	rence number	II.b.					
	I, the unde	ersigned, of	fficial veterinarian, hereb	y certify th	nat:							
	II.1.	The expo	rting country									
					exporting countr							
Part II: Certification					, , ,	, , , ,						
		II.1.1. has been free from rinderpest, peste des petits ruminants, sheep and goat pox, contagious pleuropneumonia, and Rift Valley fever during the 12 months immediately prior to collection ova (¹)/embryos (¹) to be exported and until their date of dispatch to the Union and no vaccination these diseases took place during that period; ter [II.1.2] has been free from foot-and-mouth disease during the 12 months immediately prior to collection										
	(¹) either	[11.1.2.		s been free from foot-and-mouth disease during the 12 months immediately prior to collection of t a (¹)/embryos (¹) and did not carry out vaccination against foot-and-mouth disease during that period s not been free from foot-and-mouth disease during the 12 months immediately prior to collection								
	(¹) or	[II.1.2.	the ova (¹)/embryos (¹) and the donor females disease during 30 days foot-and-mouth diseas	not been free from foot-and-mouth disease during the 12 months immediately prior to collection of ova (¹)/embryos (¹) and/or carried out vaccination against foot-and-mouth disease during that period the donor females come from holdings on which no animal was vaccinated against foot-and-mouth ase during 30 days prior to collection and no animal of susceptible species showed clinical signs of and-mouth disease during the 30 days prior to, and at least 30 days after, the ova (¹)/embryos (¹)								
	 II.2. The ova (¹)/embryos (¹) to be exported: 											
		II.2.1.	were collected (¹)/prod incidence of foot-and-r prior to their collection;	mouth dis								
		II.2.2.	were stored at all times foot-and-mouth diseas 30 days thereafter;									
		II.2.3.	were collected (¹)/prod supervised in accordar and embryo production	nce with th	ne conditions for	the approval and su	pervision of en	nbryo collection team				
		II.2.4.	meet the conditions for	ova and	embryos laid dov	wn in Chapter III(II)	of Annex D to [Directive 92/65/EEC;				
		II.2.5.	come from the donor fe	emales of	ovine (1)/caprine	(¹) species which:						
	(¹) either	[II.2.5.1.	were kept in a blueton the ova (¹)/embryos (¹)		free country or z	one for at least 60	days prior to, a	nd during collection				
	(¹) or	[11.2.5.1.	were kept during a blue	etongue v	irus seasonally f	ree period in a seas	onally free zon	e;]				
	(¹) or	[II.2.5.1.	were kept protected f ova (¹)/embryos (¹);]	the collection of th								
	(¹) or	[II.2.5.1.	underwent a serological test for the detection of antibody to the bluetongue virus serogr in accordance with the Manual of Diagnostic Tests and Vaccines for Terrestrial Animals I 60 days after collection of the ova (¹)/embryos (¹) and giving negative results.]									
	(¹) or	[II.2.5.1.	underwent an agent id Diagnostic Tests and ova (¹)/embryos (¹) coll	Vaccines	for Terrestrial	Animals on a bloc	od sample take	en on the day of th				
		II.2.5.2.	to the best of my know holding, in which, base by the owner, any of th points (a) to (d) prior to	ed on the ne followir	official notification ng diseases has	n system and acco been clinically dete	rding to the wri cted within the	itten declaration mad				
						ts (<i>Mycoplasma a</i> ge colony'), within tl		coplasma capricolur ths;				
					and the second second		0					

(b) paratuberculosis and caseous lymphadenitis, within the last 12 months;

▼<u>M4</u>

▼	M4

COUNTRY	lealth informa	ition	II.a. Certificate reference number	Ovine and caprine ova/embry
			adenomatosis, within the last three years;	
	(¹) either		a for sheep or caprine viral arthritis/encephalitis f	or goate, within the last three years."
	(¹) or			
	() 07	and all the	a for sheep or caprine viral arthritis/encephalitis infected animals were slaughtered and rema o two tests carried out at least six months apart;]	ining animals subsequently reacte
	II.2.5.3.	showed no clinical	signs of disease on the day of the ova (¹)/embryo	os (¹) collection;
(¹)(⁴) eithe	r [II.2.5.4.	originate from the (<i>B. melitensis</i>)-free	region described in Box I.8., which has been , and]	recognised as officially brucellosi
(¹) or	[11.2.5.4.		a holding which has obtained and maintained its rdance with Directive 91/68/EEC, and]	s officially brucellosis (B. melitensis,
(¹) or	[II.2.5.4.	been free from an and caprine anima Rev. 1 vaccine mo have been subjecto 	olding, where in respect of brucellosis (<i>B. meli</i> y clinical or any signs of this disease for the als have been vaccinated against this dise re than two years ago, and all ovine and capr ed to at least two tests (³), carried out with ne fate) and on	last 12 months, none of the ovin ease, save those vaccinated wit ine animals over six months of ag agative results on samples taken o
and		have not been kept	t previously in a holding of a lower status;	
(¹) either	[11.2.5.5.	have remained in ova (¹)/embryos (¹)	the exporting country for at least the past si to be exported;]	ix months prior to collection of th
(¹) or	[11.2.5.5.	health conditions a and they have be	r months prior to collection of the ova (¹)/embryo applying to donors of the ova/embryos ⁽¹⁾ which a en imported into the exporting country at leas from	are intended for export to the Unio
	II.2.5.6.	have been kept cor	ntinuously since birth in a country where the follo	wing conditions are fulfilled:
	II.2.5.6.1.	classical scrapie is	compulsorily notifiable;	
	II.2.5.6.2.	an awareness, surv	veillance and monitoring system is in place;	
	II.2.5.6.3.	ovine and caprine a	animals affected with classical scrapie are killed a	and completely destroyed;
	II.2.5.6.4.		ne and caprine animals of meat-and-bone mea ffectively enforced in the whole country for a per	
(¹) either	[11.2.5.7.	in a holding or hold the embryos to be	ntinuously for the last three years before the coll dings which has/have been complying for the las exported with the requirements laid down in poir nex VIII to Regulation (EC) No 999/2001;]	at three years before the collection of
(¹) or	[11.2.5.7.	are ovine animals a	and the embryos	
	(¹) either	[are of the ARR/AR	R prion protein genotype;]]	
	(¹) or	[carry at least one /	ARR allele and were collected after the date of 1	January 2015;]]
	[11.2.6.	were collected (1)/p	produced (¹) in the exporting country,	
(¹) either	[II.2.6.1.	which according to	official findings is free from epizootic haemorrha	gic disease (EHD);]]
(¹)(⁵) or	[11.2.6.1.	exist:a	to official findings the following serotypes of ep nd the donor females of ovine (¹)/caprine (¹) s e to the following tests carried out in an approved	pecies were subjected with negativ
	(¹) either	blood taken on tw	$\stackrel{6}{})$ for the detection of antibody to the EHD virus to occasions not more than 12 months apart for this consignment of ova (¹)/embryos (¹);]]	
	(¹) or	blood taken at inte	⁶) for the detection of antibody to the EHD virus rvals of not more than 60 days throughout the c nal collection for this consignment of ova (¹)/emb	ollection period and between 21 an

▼ <u>M4</u>

II. I	Health inform	mation II.a. Certificate reference number II.b.
	(¹) or	[an agent identification test (⁶), carried out on samples of blood collected at commencement and conclusio of, and at least every 7 days, if carried out as virus isolation test, or at least every 28 days, if carried out a polymerase chain reaction, during collection for this consignment of ova (¹)/embryos (¹);]]
	II.2.7.	were collected $(^{1})$ /produced $(^{1})$ after the date on which the embryo collection team was approved by th competent authority of the exporting country;
	II.2.8.	were processed and stored under approved conditions for at least 30 days immediately after the collection $(^1)$ /production $(^1)$ and transported under conditions for ova and embryos laid down i Chapter III(II) of Annex D to Directive 92/65/EEC;
	II.2.9.	were sent to the place of loading in a sealed container in accordance with the requirements for th transport of embryos laid down in point 6 of Chapter III(II) of Annex D to Directive 92/65/EEC and bearing the number detailed in Box I.23.
	(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 <i>in vitro</i> fertilisation (¹) using semen coming from semen collection centre approved (⁷) in accordance with:
(¹) either	[II.2.10.1.	Article 11(2) of Directive 92/65/EEC and located in a Member State of the European Union; and the seme complies with the requirements of Directive 92/65/EEC.]]
(¹) or	[II.2.10.1.	Article 17(3)(b) of Directive 92/65/EEC and located in a third country or part thereof listed in Annex I to Decision 2010/472/EU, and the semen complies with the requirements set out in Part 2 of Annex II to the Decision.]]
Notes		
Part I:		
Box I.6:	Person r	responsible for the load in EU: this box is to be filled in only if it is a certificate for transit commodity.
Box I.11:	ova/emb	origin shall correspond to the approved embryo collection team or embryo production team by which the ryos were collected/produced, processed and stored; and listed in accordance with Article 17(3)(b) or 92/65/EEC on the Commission website: http://ec.europa.eu/food/animal/semen_ova/ovine/index_en.htm.
Box I.22:	Number	of packages shall correspond to the number of containers.
Box I.23:	Identifica	ation of container and seal number shall be indicated.
Box I.26:	Fill in ac	cording to whether it is a transit or an import certificate.
Box I.27:	Fill in ac	cording to whether it is a transit or an import certificate.
Box I.28:	Species:	select amongst 'Ovis aries' or 'Capra hircus' as appropriate.
	<i>Categor</i> y embryos	y: specify if <i>in vivo</i> derived embryos, <i>in vivo</i> derived ova, <i>in vitro</i> produced embryos or micromanipulate
	Donor id	lentity shall correspond to the official identification of the animal.
	Date of c	collection shall be indicated for in vivo derived embryos and in the following format: dd.mm.yyyy.
	Date of f	freezing shall be indicated in the following format: dd.mm.yyyy.
	production and lister	I number of the team: shall correspond to the approved embryo collection team or embryo on team by which the ova/embryos were collected/produced, processed and storec d in accordance with Article 17(3)(b) of Directive 92/65/EEC on the Commission website europa.eu/food/animal/semen_ova/ovine/index_en.htm.
Part II:		
	te as appro	nriate
() Dele	as appro	priate.

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cou	NTRY			Ovine and caprine ova/embryos						
II.	Health information	II.a.	Certificate reference number	II.b.						
(³)	Tests shall be carried out in accord	dance with Anr	nex C to Directive 91/68/EEC.							
(4)	Only for the territory appearing No 206/2010 (OJ L 73, 20.3.2010,		'V' in column 6 of Part 1 of Anne	ex I to Commission Regulation (EU)						
(⁵)	See remarks for exporting country	or part thereof	concerned in Annex III to Decision 20	010/472/EU.						
(⁶)	Standards for EHD virus diagnos Vaccines for Terrestrial Animals.	stic tests are	described in Chapter 2.1.3. of the 0	OIE Manual of Diagnostic Tests and						
(7)	Only approved semen collection on the Commission websites:	entres listed in	n accordance with Article 11(4) and A	Article 17(3)(b) of Directive 92/65/EEC						
	http://ec.europa.eu/food/animal/ap http://ec.europa.eu/food/animal/se		ishments/establishments_vet_field_er e/index_en.htm.	ı.htm;						
_	The signature and the stamp must	be in a differe	nt colour to that of the printing.							
Offi	cial veterinarian									
	Name (in capital letters): Qualification and title:									
	Date:	Date: Signature:								
	Stamp:									