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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:

<u>B</u>

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
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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)

THE EUROPEAN COMMISSION,

Having regard to the Treaty on the Functioning of the European Union,

Having regard to Council Directive 92/65/EEC of 13 July 1992 laying down animal health requirements governing trade in and imports into the Community of animals, semen, ova and embryos not subject to animal health requirements laid down in specific Community rules referred to in Annex A(I) to Directive 90/425/EEC (¹), and in particular Article 17(2)(b), Article 17(3), the first indent of Article 18(1), and the introductory phrase and point (b) of Article 19 thereof,

Whereas:

- (1) Directive 92/65/EEC lays down the animal health conditions governing imports into the Union of semen, ova and embryos of animals of the ovine and caprine species ('the commodities'). It provides only commodities that come from a third country included on a list of third countries drawn up in accordance with that Directive and accompanied by a health certificate corresponding to a model also drawn up in accordance with that Directive, may be imported into the Union. The health certificate must certify that commodities come from approved collection and storage centres or collection and production teams offering guarantees at least equivalent to those laid down in Annex D(I) to that Directive.
- (2) Commission Decision 2008/635/EC of 22 July 2008 on imports of semen, ova and embryos of the ovine and caprine species into the Community as regards lists of third countries and of semen collection centres and embryo collection teams, and certification requirements (²) currently sets out the list of third countries from which Member States are to authorise imports of the commodities.
- (3) Directive 92/65/EEC, as amended by Council Directive 2008/73/EC (3), introduced a simplified procedure for the listing of semen collection and storage centres and embryo collection and production teams in third countries approved for imports of the commodities into the Union.
- (4) In addition, Annex D to Directive 92/65/EEC, as amended by Commission Regulation (EU) No 176/2010 (4), sets out certain new requirements for the commodities which are to apply from 1 September 2010. It introduces rules concerning semen storage centres and detailed conditions for their approval and supervision. It also sets out detailed conditions for the approval and supervision of embryo collection and production teams, for the collection and processing of *in vivo* derived embryos and the

⁽¹⁾ OJ L 268, 14.9.1992, p. 54.

⁽²⁾ OJ L 206, 2.8.2008, p. 17.

⁽³⁾ OJ L 219, 14.8.2008, p. 40.

⁽⁴⁾ OJ L 52, 3.3.2010, p. 14.

production and processing of *in vitro* fertilised embryos and micromanipulated embryos. It also amended the conditions to be applied to the donor animals of semen, ova and embryos of animals of the ovine and caprine species.

- (5) Accordingly, it is necessary to establish new health certificates for imports into the Union of the commodities taking into account the amendments made to Directive 92/65/EEC by Directive 2008/73/EC and Regulation (EU) No 176/2010.
- In addition, it is appropriate that consignments of the (6) commodities imported into the Union from Switzerland are accompanied by a health certificate drawn up in accordance with the models used for trade within the Union in semen, ova and embryos of animals of the ovine and caprine species set out in Commission Decision 2010/470/EU of 26 August 2010 laying down model health certificates for trade within the Union of semen, ova and embryos of animals of the equine, ovine and caprine species and in ova and embryos of animals of the porcine species (1), with the adaptations set out in point 7 of Chapter IX(B) of Appendix 2 of Annex 11 to 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 Confederation (2).
- (7) In the application of this Decision, account should be taken of the specific certification requirements and model health attestations which may be laid down in accordance with the Agreement between the European Community and the Government of Canada on sanitary measures to protect public and animal health in respect of trade in live animals and animal products (3), as approved by Council Decision 1999/201/EC (4).
- (8) In the application of this Decision, account should also be taken of the specific certification requirements and model health attestations which may be laid down in accordance with the Agreement between the European Community and New Zealand on sanitary measures applicable to trade in live animals and animal products (5), as approved by Council Decision 97/132/EC (6).
- (9) In the interest of clarity and consistency of Union's legislation, Decision 2008/635/EC should be repealed and replaced by this Decision.
- (10) To avoid any disruption of trade, the use of health certificates issued in accordance with Decision 2008/635/EC should be authorised during a transitional period subject to certain consitions.

⁽¹⁾ See page 15 of this Official Journal.

⁽²⁾ OJ L 114, 30.4.2002, p. 1.

⁽³⁾ OJ L 71, 18.3.1999, p. 3. (4) OJ L 71, 18.3.1999, p. 1.

⁽⁵⁾ OJ L 57, 26.2.1997, p. 5.

⁽⁶⁾ OJ L 57, 26.2.1997, p. 3.

(11) The measures provided for in this Decision are in accordance with the opinion of the Standing Committee on the Food Chain and Animal Health,

HAS ADOPTED THIS DECISION:

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

This Decision shall apply from 1 September 2010.

▼<u>B</u>

Article 8

Addressees

This Decision is addressed to the Member States.

▼<u>M1</u>

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 (if appropriate)	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.		
CA	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				
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).

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.
- 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.

▼<u>B</u>

PART 2

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

▼<u>M3</u>

Section A

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

COL	JNTR'	Υ			Veterinary certificate to EU		
	1.1.	Consignor Name	I.2. Certifical	te reference No	1.2.a.		
		Address Tel.	I.3. Central	competent author	ity		
ŧ			I.4. Local co	mpetent authority	1		
dispatched consignment	1.5.	Consignee Name Address Postal code	I.6. Person r Name Address	responsible for th	e load in EU		
tche		Tel.	Tel.	ode			
	1.7.	Country of origin ISO code I.8. Region of origin Code	I.9. Country destinati		de I.10. Region of Code destination		
ails	1.11.	Place of origin	I.12. Place of	destination			
Part I: Details of		Name Approval number Address	Name Address				
Parl		Name Approval number Address	Postal c	ode			
		Name Approval number Address					
	1.13.	Place of loading	I.14. Date of departure				
	1.15.	Means of transport	I.16. Entry BIP in EU				
		Aeroplane					
		Road vehicle Other	1.17.				
		Identification Documentary references					
	1.18.	Description of commodity		L19 Commodit	ty code (HS code)		
				1.10. Commodit	05 11 99 85		
					I.20. Quantity		
	1.21.				I.22. Number of packages		
	1.23.	Seal/container No			1.24.		
	1.25.	Commodities certified for:			Jac ^e		
		Artificial reproduction					
	1.26.	For transit through EU to third country	I.27. For impo	ort or admission i	into EU		
		Third country ISO code					
	1.28.	Identification of the commodities					
		Species Breed Donor identity (Scientific name)	Date of collectic		umber of the Quantity untre		

COUNTRY			Ov	rine and caprine semen — Section A			
II.	Health in	nformation II.a. C	ertificate reference No	II.b.			
	I, the un	ndersigned, official veterinarian, hereby certify that:					
II.1. The exporting country (name of exporting country) (2)							
<u> </u>	ontagious caprine pleuropneumonia and en to be exported and until its date of ng that period;						
	II.1.2.	has been free from foot-and-mouth disease during the exported and until its date of dispatch to the Union and					
II.1.2. has been free from foot-and-mouth disease during the 12 months immediately prior to collect exported and until its date of dispatch to the Union and no vaccination against this disease took II.2. The semen collection centre described in Box I.11 and at which the semen to be exported was collected.							
3	II.2.1.	II.2.1. meets the conditions for the approval of semen collection centres laid down in Chapter I(I)(1) of Annex D to Directive 92/65/EEC;					
II.2.2. is operated and supervised in accordance with the conditions applicable to semen collection cerlaid down in Chapter I(II)(1) of Annex D to Directive 92/65/EEC.							
II.3.	The ovir	ne (1)/caprine (1) animals standing at the semen collection	n centre:				
	II.3.1.	prior to their stay in the quarantine accommodation des	scribed 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 (B. n							
(¹) or	or [II.3.1.1. have belonged to a holding which has obtained and maintained its officially brucellosis (<i>B. melitensis</i>)-free 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 any signs of this disease for the last 12 months, none of the ovine and caprine animals have been vaccinated this disease, save those vaccinated with Rev. 1 vaccine more than two years ago, and all ovine and caprine are six months of age have been subjected to at least two tests (3), carried out with negative results on sample						
and		have not been kept previously in a holding of a lower s	status;				
	II.3.1.2.	have been kept continuously for at least 60 days on a has been diagnosed in the last 12 months,	holding where no case of	contagious epididymitis (Brucella ovis)			
(¹) and		[they are animals of the ovine species and have und accommodation described in point II.3.3 a complemen sensitivity and specificity, to detect contagious epididyn	t fixation test, or any other	er test with an equivalent documented			
	II.3.1.3.	to the best of my knowledge do not come from holdings based on the official notification system and according diseases has been clinically detected within the periods accommodation described in point II.3.3.	to the written declaration n	nade by the owner, any of the following			
		(a) contagious agalactia of sheep or goats (Mycoplasm mycoides "large colony"), within the last six months		capricolum, Mycoplasma mycoides var.			
		(b) paratuberculosis and caseous lymphadenitis, within	the last 12 months;				
		(c) pulmonary adenomatosis, within the last three year	rs;				
	(¹) eithei	r [(d) Maedi/Visna for sheep or caprine viral arthritis/ence	ephalitis for goats, within t	the last three years;]			
	(¹) or	[(d) Maedi/Visna for sheep or caprine viral arthritis/enco animals were slaughtered and remaining animals su months apart:]					
	II.3.2.	have undergone the following tests carried out on commencement of the period of quarantine specified in		d within the 28 days preceding the			

II.	Health	information	II.a. Certificate reference No	II.b.				
		brucellosis (B. melitensis), with negative result	ts in each case in accordance with	Annex C to Directive 91/68/EEC				
		contagious epididymitis (<i>Brucella. ovis</i>), in the ca Annex D to Directive 91/68/EEC, or any other t						
		— border disease in accordance with point 1.4(c) of Chapter II(II) of Annex D to Directive 92/65/EEC;						
	II.3.3.	have satisfied the quarantine isolation period of at least 28 days in a quarantine accommodation specifically approved for to purpose by the competent authority and during that period:						
	II.3.3.1.	only animals of at least the same health status we	re present in the quarantine accomm	nodation;				
	II.3.3.2.	the animals have undergone the following tests, of the exporting country on samples taken not exaccommodation, for:						
		- brucellosis (B. melitensis) with negative results	s in each case in accordance with	Annex C to Directive 91/68/EEC				
		 contagious epididymitis (<i>Brucella ovis</i>), in the ca Annex D to Directive 91/68/EEC, or any other t 						
		— border disease in accordance with point 1.6 of	Chapter II(II) of Annex D to Directive	92/65/EEC;				
	II.3.4. have undergone at least once a year the routine tests for: — brucellosis (B. melitensis) with negative results in each case in accordance with Annex C to Directive							
		 contagious epididymitis (<i>Brucella ovis</i>), in the ca Annex D to Directive 91/68/EEC, or any other to 						
		- border disease in accordance with point 5(c) of	f Chapter II(II) of Annex D to Directive	re 92/65/EEC.				
11.4.	The ser	men to be exported was obtained from donor rams ((1)/bucks (1) which:					
	II.4.1.	were admitted to the approved semen collection of	entre with the express permission of	the centre veterinarian;				
	II.4.2.	show no clinical signs of disease on the day of a semen was collected;	dmission to the approved semen co	ellection centre and on the day the				
(¹) either	[11.4.3.	have not been vaccinated against foot-and-mouth of	disease during the 12 months prior t	o collection of the semen;]				
(1) or	[11.4.3.	have been vaccinated against foot-and-mouth disea five straws) of each collection have been submitted						
	II.4.4.	have been kept at an approved semen collection collection of the semen, in the case of collections		least 30 days immediately prior to				
	II.4.5.	have not served naturally after their entry to the quathe day of semen collection;	arantine accommodation described in	point II.3.3 and up to and including				
	II.4.6.	have been kept at approved semen collection cent	tres:					
	II.4.6.1.	which have been free from foot-and-mouth disease after collection or, in the case of fresh semen, until t kilometres radius in which there has been no case semen;	the date of dispatch, and which are sit	tuated in the centre of an area of 10				
	II.4.6.2.	which have been free, during the period commenci semen or, in the case of fresh semen, until the da (Bruceila. ovis), anthrax and rables;						

COUNTRY			Ovine	and caprine semen — Section A			
II.	Health info	ormation	II.a. Certificate reference No	II.b.			
(¹) either	[11.4.7.	have remained in the exporting country f exported;]	for at least the past six months prior	to collection of the semen to be			
(¹) or	[11.4.7.	[II.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	[II.4.8.	were kept in a bluetongue virus-free country	y or zone for at least 60 days prior to, a	and during, collection of the semen;]			
(¹) or	[II.4.8.	were kept during a bluetongue virus seasor during collection of the semen;]	nally free period in a seasonally free zo	one for at least 60 days prior to, and			
(†) or	[11.4.8.	were kept in a vector-protected establishr	ment for at least 60 days prior to, ar	nd during collection of the semen;]			
(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 final 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 seven 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 exporting country whic (EHD);]	h according to official findings is free fi	rom epizootic haemorrhagic disease			
(¹) or	[11.4.9.	were resident in the exporting country in haemorrhagic disease (EHD) exist:each case to:					
	(¹) either	[a serological test (6) for the detection of samples of blood taken on two occasions the final collection for this consignment of	not more than 12 months apart prior				
	(¹) or	[a serological test (6) for the detection of samples of blood taken at intervals of not n days after the final collection for this consi	nore than 60 days throughout the collec				
	(¹) or	[an agent identification test (⁶) carried out in conclusion of, and at least every seven day for this consignment of semen.]]					
	II.4.10.	have been kept continuously since birth in	a country where the following condition	ons are fulfilled:			
	II.4.10.1.	classical scrapie is compulsorily notifiable;					
	II.4.10.2.	an awareness, surveillance and monitoring	system is in place;				
	II.4.10.3.	ovine and caprine animals affected with cla	assical scraple are killed and complete	ely destroyed;			
	II.4.10.4.	the feeding to ovine and caprine animals of effectively enforced in the whole country for					
(¹) either	[II.4.11.	have been kept continuously for the last thr holdings which has/have been complying f with the requirements laid down in points 1 999/2001;]	or the last three years before the colle	ection of the semen to be exported			
(1) or	[II.4.11.	are ovine animals of ARR/ARR prion prote	ain genotype.]				

COUNTRY			Ovine	and caprine semen — Section A		
II.	Health inf	ormation	II.a. Certificate reference No	II.b.		
11.5.	The seme	en to be exported:				
	II.5.1.	was collected after the date on which the exporting country;	semen collection centre was approved	d by the competent authority of the		
	II.5.2.	was collected, processed, preserved, sto semen laid down in Chapter III(I) of Annex		vith the requirements applicable to		
	II.5.3.	was sent to the place of loading in a sealer trade laid down in point 1.4 of Chapter III(I Box I.23.				
(1) other	[II.6.	No antibiotics were added to the semen.]				
(¹) or	[II.6.	The following antibiotic or combination of a of not less than (7) :	ntibiotics was added to produce a con	centration in the final diluted semen		
				.]		
Notes						
Part I:						
Box 1.6: F	erson respons	tible for the load in EU: this box is to be filled	in only if it is a certificate for transit of	commodity.		
		shall correspond to the approved semen collect of Directive 92/65/EEC on the Commission w				
Box 1.22: N	lumber of pack	kages shall correspond to the number of conta	ainers.			
Box 1.23: 10	dentification of	container and seal number shall be indicated				
Box 1.26: F	ill in according	to whether it is a transit or an import certification	ate.			
Box 1.27: F	ill in according	to whether it is a transit or an import certification	ate.			
Box 1.28: S	pecies: select	amongst "Ovis aries" or "Capra hircus" as ap	oropriate.			
[onor identity s	shall correspond to the official identification of	the animal.			
[ate of collection	on shall be indicated in the following format: d	d.mm.yyyy.			
A	pproval numbe	er of the centre shall correspond to the appro-	val number of the semen collection ce	entre indicated in Box I.11.		
Part II:						
(1) Delete a	as necessary.					
(2) Only thi	rd countries lis	sted in Annex I to Decision 2010/472/EU.				
(3) Tests s	nall be carried	out in accordance with Annex C to Directive	91/68/EEC.			
	(4) Only for the territory appearing with the entry "V" in column 6 of Part 1 of Annex I to Commission Regulation (EU) No 206/2010 (OJ L 73, 20.3.2010, p. 1).					
(5) See ren	narks for expo	rting country concerned in Annex I to Decision	2010/472/EU.			
(⁶) Standar Animals		rus diagnostic tests are described in Chapter 2	2.1.3. of the OIE Manual of Diagnostic	e Tests and Vaccines for Terrestrial		
(7) Insert n	ames and con-	centrations.				
— The sig	nature and the	stamp must be in a different colour to that of	the printing.			

COU	INTRY	Ovine and caprine semen — Section A
II.	Health information	II.a. Certificate reference No II.b.
Offi	cial veterinarian	-
	Name (in capital letters):	Qualification and title:
	Date:	Signature:
	Stamp:	

Section B

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

cou	COUNTRY:										Veterinary cer	tificate to EU
	1.1.	Consignor Name				1.2.	Certificate	referenc	e No	1.	2.a.	
		Address Tel. Consignee					Central o	ompetent	authority	′		
							Local cor	npetent a	uthority			
	I.5.						Person re	sponsible	for the	load in E	U	
ent		Name					Name					
gnm		Address					Address					
hed consi		Postal code Tel.					Postal cor Tel.	de				
Part I: Details of dispached consignment	1.7.	Country of origin	ISO code	I.8. Region of origin	Code	1.9.	Country of destination	of n	ISO code	l.10.	Region of destination	Code
Details	1.11.	Place of origin				l.12.	Place of o	destination	n			
벌		Name		Approval number			Name					
ă.		Address		A			Address					
		Name Address		Approval number			Postal co	46				
		Name		Approval number			i Ustai CO	ac .				
		Address										
	I.13.	I.13. Place of loading			1.14.	I.14. Date of departure						
	I.15. Means of transport			1.16.	Entry BIP	in EU						
		Aeroplane	Ship [☐ Railway wa	gon 🔲							
		Road vehicle	Other			1.17.	I.17. No(s) of related original certificates					
		Identification										
	1.18.	Documentary refer Description of com						I.19. Cor	nmodity i	code (HS	code)	
		'	,							05 11 9		
									1.2	0. Quan	tity	
	I.21.								1.2	2. Numb	er of packages	
	1.23.	Seal/container No							1.2	4.		
	1.25.	Commodities certif	ied for:									
		Artificial reproducti	on									
	1.26.	For transit through	the EU to a	third country		1.27.	For impor	t or admis	ssion into	the EU		
		Third country		ISO code								
	1.28.	Identification of the	commoditie	es .								
		Species (scientific name)	В	reed Donor identi	ty Da	te of co	lection			al numbo e centre	er	Quantity
	1											

	COUNTRY	/ :			Ovine	and caprine semen — Section A	
	II. H	-lealth	information		II.a. Certificate reference No	II.b.	
	I, the und	dersign	ed official ve	terinarian of		hereby certify that:	
				(name c	of exporting country) (²)		
			II.1.	The centre (3) described in Box I.11 at whi	ch the semen to be exported to the E	European Union was stored:	
	(1) either		[11.1.1.	meets the conditions laid down in Chapter	I(I)(1) of Annex D to Directive 92/65/I	EEC;	
and II.1.2. is operated and supervised in accordance with the conditions laid down in Chapter I(II)(1) of Annex D to 92/65/EEC.] iii meets the conditions laid down in Chapter I(I)(2) of Annex D to Directive 92/65/EEC; and II.1.2. is operated and supervised in accordance with the conditions laid down in Chapter I(II)(2) of Annex D to Directive 92/65/EEC;					pter I(II)(1) of Annex D to Directive		
					EEC;		
Part	and II.1.2. is operated and supervised in accordance with the conditions laid down in Chapter I(II)(2) of Annex D to I 92/65/EEC.]					pter I(II)(2) of Annex D to Directive	
			II.2.	The semen to be exported to the European	n Union:		
	-		II.2.1.	has been collected, processed and stored approved semen collection centre (*) opera of Annex D to Directive 92/65/EEC, and			
			(¹) either	[located in the exporting country;]			
			(1) and/or	[located in(5);			
			and	has been imported to the exporting country caprine species into the European Union in			
			II.2.2.	was moved to the centre described in Part I to Decision 2010/472/EU (6):]	t I.11 under conditions at least as strict as in Section A of Part 2 of Annex II		
			II.2.3.	was stored under conditions which satisfy	the terms of Annex D to Directive 92/	'65/EEC;	
			II.2.4.	was sent to the place of loading in a seale Directive 92/65/EEC and bearing the numb	aled container in accordance with point 1.4 of Chapter III(I) of Annex D to ober indicated in Box I.23.		
	Notes						
	Part I:						
	Box I.11:	place	of origin sha	all correspond to the approved semen storage	ge centre of dispatch of the semen.		
	Box 1.17:	above	e from the ap	o the serial number of the individual official de proved semen collection centre of its origin to or the officially endorsed copies of thereof	o the centre described in Box I.11. The		
	Box 1.22:	numb	er of packag	es shall correspond to the number of contain	ners.		
	Box 1.23:	identi	fication of co	ntainer and seal number shall be indicated.			
	Box 1.28:	donor	r identity shal	Il correspond to the official identification of the	he animal.		
		date	of collection s	shall be indicated in the following format: dd	l/mm/yyyy.		
		appro		of the centre shall correspond to the approval	number of the approved semen collect	ction centre in which the semen was	

Ovine and caprine semen — Section B

COUNTRY:

II.	Health Information	II.a. Certificate reference No	II.b.					
Pai	Part II:							
(1)	(1) Delete as necessary.							
(2)	(2) Only third countries listed in Annex I to Decision 2010/472/EU.							
(3)	(3) Only approved semen collection or storage centres listed in accordance with Article 17(3)(b) of Directive 92/65/EEC on the Commission website:							
	http://ec.europa.eu/food/animal/semen_ova/ovine/index_en.htm							
(4)	4) Only approved semen collection centres listed in accordance with Article 11(4) and 17(3)(b) of Directive 92/65/EEC on the Commission websites:							
	http://ec.europa.eu/food/animal/approved_establishments/establishme http://ec.europa.eu/food/animal/semen_ova/ovine/index_en.htm	nts_vet_field_en.htm						
(5)	Only third countries listed in Annex I to Decision 2010/472/EU and to	he EU Member States.						
(⁶)	(6) The original(s) of the document(s) or the health certificate(s) or the officially endorsed copies of thereof that accompanied the semen described above from the approved semen collection centre in which the semen was collected to the approved semen storage centre of the semen dispatch described in Box I.11 must be attached to this certificate.							
Off	icial veterinarian (*)							
	Name (in capital letters): Qualification and title:							
	Date:	Si	gnature:					
	Stamp:							
(*)	(*) The signature and the stamp must be in a different colour to that of the printing.							

▼<u>M1</u>

▼<u>M2</u>

▼<u>M1</u>

ANNEX III

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

		Remarks			
ISO Code	Name of the third country	Description of the territory (if appropriate)	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.		
CA	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				
IS	Iceland				
NZ	New Zealand				
PM	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.		

⁽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.

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.
- 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 certificate for imports of consignments of ova and embryos of animals of the ovine and caprine species

OUN	TRY:			Veterinary certificate to EL					
	I.1. Consignor Name			Certificate reference No I.2.a.					
		Address	I.3. Central competent authority						
		Tel.	I.4. Local competent authority						
ignment	1.5.	Consignee Name Address		I.6. Person responsible for the load in EU Name Address					
Part I: Details of dispatched consignment		Postal code Tel.	Postal code Tel.						
of dispat	1.7.	Country of ISO code I.8. Region of Code origin origin	1.9.	Country of ISO code I.10. Region of Code destination destination					
tails	l.11.	Place of origin	l.12.	Place of destination					
Part I: De		Name Approval number Address Name Approval number Address Name Approval number Address		Name Address Postal code					
	I.13.	Place of loading	l.14.	Date of departure					
	l.15.	Means of transport	I.16.	Entry BIP in EU					
		Aeroplane Ship Railway wagon Road vehicle Other Identification Documentary references	l.17.						
	1.18.	Description of commodity		I.19. Commodity code (HS code) 05 11 99 85					
				I.20. Quantity					
	l.21.			I.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 1.27. For import or admission into EU							
		Third country ISO code							
	1.28.	Identification of the commodities							
	(Sc	Species Category Donor identity Date of collect cientific name)	tion [Date of freezing Approval number of the team Quantity					

COUNTRY	Health infor	mation II.a. Certificate reference n	Ovine and caprine ova/embryo umber II.b.					
		ficial veterinarian, hereby certify that:						
,								
II.1.	The exporting country							
	(name of exporting country) (²)							
	II.1.1.	has been free from rinderpest, peste des petits ruminants, sheep and goat pox, contagious capi pleuropneumonia, and Rift Valley fever during the 12 months immediately prior to collection of ova (')/embryos (') to be exported and until their date of dispatch to the Union and no vaccination aga these diseases took place during that period;						
(1) either	[II.1.2.	has been free from foot-and-mouth disease during the 12 months immediately prior to collection of the ova (¹)/embryos (¹) and did not carry out vaccination against foot-and-mouth disease during that period;]						
(¹) or	[II.1.2.	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;]						
11.2.	The ova (The ova (¹)/embryos (¹) to be exported:						
	II.2.1.	2.1. were collected (¹)/produced (¹) and processed on premises within a 10-km radius of which there was r incidence of foot-and-mouth disease, vesicular stomatitis, Rift Valley fever in the 30 days immediate prior to their collection;						
	II.2.2.	were stored at all times on approved premises within a foot-and-mouth disease, vesicular stomatitis or Rift Va 30 days thereafter;						
	II.2.3.	were collected (¹)/produced (¹) by the team described supervised in accordance with the conditions for the app and embryo production teams laid down in Chapter I(III)	roval and supervision of embryo collection team					
	II.2.4.	meet the conditions for ova and embryos laid down in Ch	napter III(II) of Annex D to Directive 92/65/EEC;					
	II.2.5.	come from the donor females of ovine (1)/caprine (1) spec	cies which:					
(1) either	[II.2.5.1.	were kept in a bluetongue virus-free country or zone for at least 60 days prior to, and during colle the ova $(^1)$ /embryos $(^1)$;]						
(1) or	[11.2.5.1.	were kept during a bluetongue virus seasonally free period in a seasonally free zone;]						
(¹) or [II.2.5.1. were kept protected from the vector for at least 60 days prior to, and during the colova (¹)/embryos (¹);] (¹) or [II.2.5.1. underwent a serological test for the detection of antibody to the bluetongue virus serogrous in accordance with the Manual of Diagnostic Tests and Vaccines for Terrestrial Animals be 60 days after collection of the ova (¹)/embryos (¹) and giving negative results;]								
					(¹) or	[II.2.5.1.	underwent an agent identification test for bluetongue virus, carried out in accordance with the M Diagnostic Tests and Vaccines for Terrestrial Animals on a blood sample taken on the day ova (¹)/embryos (¹) collection or the day of slaughtering and giving negative results;]	
	II.2.5.2.	to the best of my knowledge do not come from holdings holding, in which, based on the official notification system by the owner, any of the following diseases has been clipoints (a) to (d) prior to collection of the ova (1)/embryos	m and according to the written declaration mad inically detected within the periods referred to					
		(a) contagious agalactia of sheep or goats (Myconflasma mycoides var. mycoides 'large color						
		(b) paratuberculosis and caseous lymphadenitis, with	hin the last 12 months;					

COUNTRY		Ovine and caprine ova/embryos					
II. F	lealth informa	tion II.a. Certificate reference number II.b.					
		(c) pulmonary adenomatosis, within the last three years;					
	(1) either	[(d) Maedi/Visna for sheep or caprine viral arthritis/encephalitis for goats, within the last three years;]					
	(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 negatively to two tests carried out at least six months apart;]					
	II.2.5.3.	showed no clinical signs of disease on the day of the ova (1)/embryos (1) collection;					
(1)(4) eithei	[II.2.5.4.	originate from the region described in Box I.8., which has been recognised as officially brucellosis (<i>B. melitensis</i>)-free, and]					
(¹) or	[11.2.5.4.	have belonged to a holding which has obtained and maintained its officially brucellosis (B. melitensistere status in accordance with Directive 91/68/EEC, and]					
(¹) or	[II.2.5.4.	originate from a holding, where in respect of brucellosis (<i>B. melitensis</i>) 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					
and		have not been kept previously in a holding of a lower status;					
(¹) either	[II.2.5.5.	have remained in the exporting country for at least the past six months prior to collection of the ova $\binom{1}{2}$ -mbryos $\binom{1}{1}$ to be exported;]					
(¹) or	[II.2.5.5.	during the past six months prior to collection of the ova $\binom{1}{}$ /embryos $\binom{1}{}$ they complied with the anima health conditions applying to donors of the ova/embryos $\binom{1}{}$ which are intended for export to the Unior and they have been imported into the exporting country at least 30 days prior to collection of the ova $\binom{1}{}$ /embryos $\binom{1}{}$ from					
	II.2.5.6.	have been kept continuously since birth in a country where the following conditions are fulfilled:					
	II.2.5.6.1.	classical scrapie is compulsorily notifiable;					
	II.2.5.6.2.	an awareness, surveillance and monitoring system is in place;					
	II.2.5.6.3.	ovine and caprine animals affected with classical scrapie are killed and completely destroyed;					
	II.2.5.6.4.	the feeding to ovine and caprine animals of meat-and-bone meal, or greaves of ruminant origin has been banned and effectively enforced in the whole country for a period of at least the last seven years;					
(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 complying 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 VIII to Regulation (EC) No 999/2001;]					
(1) or	[11.2.5.7.	are ovine animals and the embryos					
	(1) either	[are of the ARR/ARR prion protein genotype;]]					
	(1) or	[carry at least one ARR allele and were collected after the date of 1 January 2015;]]					
	[II.2.6.	were collected (¹)/produced (¹) in the exporting country,					
(1) either	[II.2.6.1.	which according to official findings is free from epizootic haemorrhagic disease (EHD);]]					
(¹)(⁵) or	[II.2.6.1.	in which according to official findings the following serotypes of epizootic haemorrhagic disease (EHD exist:					
	(1) either	[a serological test $(^6)$ for the detection of antibody to the EHD virus serogroup, carried out on samples oblood taken on two occasions not more than 12 months apart prior to and not less than 21 days following collection for this consignment of ova $(^1)$ /embryos $(^1)$;]]					
	(¹) or	[a serological test (⁶) for the detection of antibody to the EHD virus serogroup, carried out 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 ova (¹)/embryos (¹);]]					

COUNTRY	•		,			Ovine and caprine ova/embryos		
II.	Health inforr	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 conclusion, and at least every 7 days, if carried out as virus isolation test, or at least every 28 days, if carried out polymerase chain reaction, during collection for this consignment of ova (¹)/embryos (¹);[]						
	II.2.7.	were collected (1)/producompetent authority of t			yo colle	ction team was approved by the		
	II.2.8.		(1) and	transported under conditions		30 days immediately after their va and embryos laid down in		
	II.2.9.		id down ir			e with the requirements for the Directive 92/65/EEC and bearing		
	(¹) [II.2.10.		sult of in			rhich were conceived by artificial ag from semen collection centres		
(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 semen complies with the requirements of Directive 92/65/EEC.]]						
(¹) or	[II.2.10.1.	I.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	esponsible for the load in	<i>EU</i> : this	box is to be filled in only if it is a c	ertificat	e for transit commodity.		
Box I.11:	ova/emb	Place 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 Article 17(3)(b) of Directive 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 corres	pond to th	e number of containers.				
Box I.23:	Identifica	ation of container and sea	al 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 an	es' or 'Ca	pra hircus' as appropriate.				
		Category: specify if in vivo derived embryos, in vivo derived ova, in vitro produced embryos or micromanipulated embryos.						
	Donor identity shall correspond to the official identification of the animal.							
	Date of c	Date of collection shall be indicated for in vivo derived embryos and in the following format: dd.mm.yyyy.						
	Date of freezing shall be indicated in the following format: dd.mm.yyyy.							
	production and liste		he ova cle 17(3	a/embryos were collected/pr 3)(b) of Directive 92/65/EE	roduced			
Part II:								
lah me								

- (¹) Delete as appropriate.
- $(^2)$ Only third countries or parts thereof listed in Annex I to Decision 2010/472/EU.

II.	Health information	II.a.	Certificate reference number	II.b.		
(³)	Tests shall be carried out in accor	dance with Ann	ex C to Directive 91/68/EEC.			
(⁴)	Only for the territory appearing with the entry 'V' in column 6 of Part 1 of Annex I to Commission Regulation (EU) No 206/2010 (OJ L 73, 20.3.2010, p. 1).					
(⁵)	See remarks for exporting country or part thereof concerned in Annex III to Decision 2010/472/EU.					
(⁶)	Standards for EHD virus diagnostic tests are described in Chapter 2.1.3. of the OIE Manual of Diagnostic Tests and Vaccines for Terrestrial Animals.					
(⁷)	Only approved semen collection centres listed in accordance with Article 11(4) and Article 17(3)(b) of Directive 92/65/EEC on the Commission websites:					
	http://ec.europa.eu/food/animal/approved_establishments/establishments_vet_field_en.htm; http://ec.europa.eu/food/animal/semen_ova/ovine/index_en.htm.					
_	The signature and the stamp must be in a different colour to that of the printing.					
Offi	cial veterinarian					
	Name (in capital letters):		Qualification and title:			
	Date:		Signature:			
	Stamp:					