

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

COUNTRY:

Veterinary certificate to EU

Part I : Details of dispatched consignment	I.1. Consignor Name Address Tel.		I.2. Certificate reference No		I.2.a.							
			I.3. Central competent authority									
			I.4. Local competent authority									
	I.5. Consignee Name Address Postal code Tel.		I.6. Person responsible for the load in EU Name Address Postal code Tel.									
			I.7. Country of origin		ISO code		I.8. Region of origin		Code			
	I.9. Country destination		of		ISO code		I.10. Region of destination		Code			
	I.11. Place of origin Name Address Name Address Name Address Approval number Approval number Approval number				I.12. Place of destination Name Address Postal code							
					I.13. Place of loading				I.14. Date of departure			
					I.15. Means of transport Aeroplane <input type="checkbox"/> Ship <input type="checkbox"/> Railway wagon <input type="checkbox"/> Road vehicle <input type="checkbox"/> Other <input type="checkbox"/> Identification Documentary references				I.16. Entry BIP in EU			
	I.18. Description of commodity				I.19. Commodity code (HS code) 05 11 99 85							
				I.20. Quantity								
I.21.				I.22. Number of packages								
I.23. Seal/Container No				I.24.								
I.25. Commodities certified for: Artificial reproduction <input type="checkbox"/>												
I.26. For transit through EU to third country <input type="checkbox"/>			I.27. For import or admission into EU <input type="checkbox"/>									
Third country			ISO code									
I.28. Identification of the commodities Species Breed Category Donor identity Date of collection Date of freezing Approval number of the team Quantity (Scientific name)												

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Ovine and caprine ova/embryos

	II. Health information	II.a. Certificate reference No	II.b.
Part II: Certification	I, the undersigned, official 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 caprine pleuropneumonia, and Rift Valley fever during the 12 months immediately prior to collection of the ova ⁽¹⁾ /embryos ⁽¹⁾ to be exported and until their date of dispatch to the Union and no vaccination against 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 <i>zona pellucida</i> ;	
	II.2.	The ova ⁽¹⁾ /embryos ⁽¹⁾ to be exported:	
		II.2.1. were collected ⁽¹⁾ /produced ⁽¹⁾ and processed on premises within a 10-km radius of which there was no incidence of foot-and-mouth disease, vesicular stomatitis, Rift Valley fever in the 30 days immediately prior to their collection;	
		II.2.2. were stored at all times on approved premises within a 10-km radius of which there was no incidence of foot-and-mouth disease, vesicular stomatitis or Rift Valley fever from the time of their collection until 30 days thereafter;	
		II.2.3. were collected ⁽¹⁾ /produced ⁽¹⁾ by the team described in Box I.11., which has been approved and supervised in accordance with the conditions for the approval and supervision of embryo collection teams and embryo production teams laid down in Chapter I(III) of Annex D to Directive 92/65/EEC;	
		II.2.4. meet the conditions for ova and embryos laid down in Chapter III(II) of Annex D to Directive 92/65/EEC;	
	II.2.5. come from the donor females of ovine ⁽¹⁾ /caprine ⁽¹⁾ species 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 collection of the ova ⁽¹⁾ /embryos ⁽¹⁾ ;		
^{(1)or}	[II.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 collection of the ova ⁽¹⁾ /embryos ⁽¹⁾ ;		
^{(1)or}	[II.2.5.1. underwent a serological test to detect antibody to the bluetongue virus group, carried out in accordance with the Manual of Diagnostic Tests and Vaccines for Terrestrial Animals between 21 and 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 Manual of Diagnostic Tests and Vaccines for Terrestrial Animals on a blood sample taken on the day of the 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 and have not been in contact with animals of a holding, in which, based on the official notification system and according to the written declaration made by the owner, any of the following diseases has been clinically detected within the periods referred to in points (a) to (d) prior to collection of the ova ⁽¹⁾ /embryos ⁽¹⁾ to be exported:		
	(a) contagious agalactia of sheep or goats (<i>Mycoplasma agalactiae</i> , <i>Mycoplasma capricolum</i> , <i>Mycoplasma mycoides var. mycoides</i> "large colony"), within the last six months;		
	(b) paratuberculosis and caseous lymphadenitis, within the last 12 months;		
	(c) pulmonary adenomatosis, within the last three years;		

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	⁽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⁾ either	[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	[II.2.5.4. have belonged to a holding which has obtained and maintained its official brucellosis (<i>B. melitensis</i>)-free status in accordance with Directive 91/68/EEC, and]		
⁽¹⁾ or	[II.2.5.4. originate from a holding, where in respect of brucellosis (<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 (date) and on (date) at least six months apart, the latter being within 30 days prior to collection of the ova ⁽¹⁾ /embryos ⁽¹⁾ .]		
<i>and</i>	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 ⁽¹⁾ /embryos ⁽¹⁾ to be exported;]		
⁽¹⁾ or	[II.2.5.5. during the past six months prior to collection of the ova ⁽¹⁾ /embryos ⁽¹⁾ they complied with the animal health conditions applying to donors of the ova/embryos ⁽¹⁾ which are intended for export to the Union and they have been imported into the exporting country at least 30 days prior to collection of the ova ⁽¹⁾ /embryos ⁽¹⁾ from ⁽²⁾ .]		
	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 1.3. (a) to (f) of Section A of Chapter A of Annex VIII to Regulation (EC) No 999/2001;]		
⁽¹⁾ or	[II.2.5.7. are ovine animals and the embryos of the ARR/ARR prion protein genotype;]		
	[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: and were subjected with negative results in each case to:		
⁽¹⁾ either	[a serological test ⁽⁶⁾ for the detection of antibody to the EHDV group carried out in an approved laboratory on samples of blood 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 EHDV group, 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⁾ .]		
⁽¹⁾ or	[an agent identification test ⁽⁶⁾ carried out in approved laboratories on samples of blood collected at commencement and conclusion of, and at least every 7 days (virus isolation test) or at least every 28 days (PCR test) during collection for this consignment of ova ⁽¹⁾ /embryos ⁽¹⁾ .]		

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Ovine and caprine ova/embryos

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	<p>II.2.7. were collected⁽¹⁾/produced⁽¹⁾ after the date on which the embryo collection team was approved by the competent authority of the exporting country;</p> <p>II.2.8. were processed and stored under approved conditions for at least 30 days immediately after their collection⁽¹⁾/production⁽¹⁾ and transported under conditions for ova and embryos laid down in Chapter III(II) of Annex D to Directive 92/65/EEC;</p> <p>II.2.9. were sent to the place of loading in a sealed container in accordance with the requirements for the 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.</p> <p>⁽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 centres approved⁽⁷⁾ in accordance with:</p> <p>⁽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.]]</p> <p>⁽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 that Decision.]]</p>		
Notes			
Part I:			
Box I.6.:	<i>Person responsible for the load in EU:</i> this box is to be filled in only if it is a certificate for transit commodity.		
Box I.11.:	<i>Place of origin</i> 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 correspond to the number of containers.		
Box I.23.:	Identification of container and seal 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.:	<p><i>Species:</i> select amongst “<i>Ovis aries</i>” or “<i>Capra hircus</i>” as appropriate.</p> <p><i>Category:</i> specify if <i>in vivo</i> derived embryos, <i>in vivo</i> derived ova, <i>in vitro</i> produced embryos or micromanipulated embryos.</p> <p><i>Donor identity</i> shall correspond to the official identification of the animal.</p> <p><i>Date of collection</i> shall be indicated for <i>in vivo</i> derived embryos and in the following format: dd.mm.yyyy.</p> <p><i>Date of freezing</i> shall be indicated in the following format: dd.mm.yyyy.</p> <p><i>Approval number of the team:</i> 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.</p>		
Part II:			
⁽¹⁾	Delete as appropriate.		
⁽²⁾	Only third countries or parts thereof listed in Annex I to Decision 2010/472/EU.		
⁽³⁾	Tests shall 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 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:		

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<p>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.</p> <ul style="list-style-type: none">The signature and the stamp must be in a different colour to that of the printing.								
<p>Official veterinarian</p> <table><tr><td>Name (in capital letters):</td><td>Qualification and title:</td></tr><tr><td>Date:</td><td>Signature:</td></tr><tr><td>Stamp:</td><td></td></tr></table>			Name (in capital letters):	Qualification and title:	Date:	Signature:	Stamp:	
Name (in capital letters):	Qualification and title:							
Date:	Signature:							
Stamp:								