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## COMMISSION DECISION

# of 2 September 1992

# concerning animal health conditions and veterinary certification for importation of bovine embryos from third countries

(92/471/EEC)

(OJ L 270, 15.9.1992, p. 27)

## Amended by:

<u>B</u>

		(	Official Jour	mal
		No	page	date
<u>M1</u>	Commission Decision 94/280/EC of 28 April 1994	L 120	52	11.5.1994
<u>M2</u>	Commission Decision 94/453/EC of 29 June 1994	L 187	11	22.7.1994
<u>M3</u>	Commission Decision 96/572/EC of 24 September 1996	L 250	20	2.10.1996
► <u>M4</u>	Commission Decision 2004/52/EC of 9 January 2004	L 10	67	16.1.2004
► <u>M5</u>	Commission Decision 2004/786/EC of 18 November 2004	L 346	32	23.11.2004
Amend	led by:			
► <u>A1</u>	Act of Accession of Austria, Sweden and Finland	C 241	21	29.8.1994
	(adapted by Council Decision 95/1/EC, Euratom, ECSC)	L 1	1	1.1.1995
► <u>A2</u>	Act concerning the conditions of accession of the Czech Republic, the Republic of Estonia, the Republic of Cyprus, the Republic of Latvia, the Republic of Lithuania, the Republic of Hungary, the Republic of Malta, the Republic of Poland, the Republic of Slovenia and the Slovak Republic and the adjustments to the Treaties on which the European Union is founded	L 236	33	23.9.2003

## **COMMISSION DECISION**

## of 2 September 1992

concerning animal health conditions and veterinary certification for importation of bovine embryos from third countries

(92/471/EEC)

THE COMMISSION OF THE EUROPEAN COMMUNITIES.

Having regard to the Treaty establishing the European Economic Community,

Having regard to Council Directive 89/556/EEC of 25 September 1989 on animal health conditions governing intra-Community trade in and importation from third countries of embryos of domestic animals of the bovine species (¹), as amended by Directive 90/425/EEC (²), and in particular Articles 9 and 10 thereof,

Whereas the list of third countries from which Member States are permitted to import bovine embryos has been established by Commission Decision 91/270/EEC (3);

Whereas Commission Decision 92/452/EEC (4) establishes the list of embryo collection teams for certain third countries; whereas Decision 92/452/EEC will be completed in due course with new information for other third countries;

Whereas it is necessary to lay down the animal health conditions and veterinary certification required for imports of bovine embryos from third countries:

Whereas the competent authorities of the third country in which the embryos intended for export to the Community were collected have undertaken to ensure that such embryos have been collected and processed by approved and supervised embryo collection teams, that they have been obtained from animals of satisfactory health status, that they have been stored and transported in accordance with the rules which preserve their health status and are accompanied during transport by an animal health certificate in order to ensure that this obligation has been fulfilled;

Whereas the competent veterinary authorities of the third countries on the list have undertaken to notify the Commission and the Member States by telex or telefax within 24 hours of the confirmation of the occurrence of any of the following diseases: rinderpest, foot-and-mouth disease, contagious bovine pleuropneumonia, bluetongue, epizootic haemorrhagic disease, Rift Valley fever and contagious vesicular stomatitis or the adoption of vaccination against them;

Whereas the animal health situation in the third countries on the list is satisfactory from the point of view of import of bovine embryos; whereas the veterinary services in these countries are well-structured and organized;

Whereas the animal health certificate is adapted to cater for the animal health situation in individual third countries;

Whereas the measures provided for in this Decision are in accordance with the opinion of the Standing Veterinary Committee,

<sup>(1)</sup> OJ No L 302, 19. 10. 1989, p. 1.

<sup>(2)</sup> OJ No L 224, 18. 8. 1990, p. 29.

<sup>(3)</sup> OJ No L 134, 29. 5. 1991, p. 56.

<sup>(4)</sup> OJ No L 250, 29. 8. 1992, p. 40.

# HAS ADOPTED THIS DECISION:

## Article 1

- 1. Member States shall authorize the importation of bovine embryos conforming to the guarantees laid down in the animal health certificate in accordance with Annex A, Part I. This certificate must accompany consignments of embryos coming from third countries or parts of third countries listed in Annex A, Part II.
- 2. Member States shall authorize the importation of bovine embryos conforming to the guarantees laid down in the animal health certificate in accordance with Annex B, Part I. This certificate must accompany consignments of embryos coming from third countries or parts of third countries listed in Annex B, Part II.

### Article 2

This Decision is addressed to the Member States.

**▼**<u>M1</u>

ANNEX A

**▼**<u>M5</u>

## Part I

VETERINARY CERTIFICATE embryos of domestic animals of the bovine species for imports collected or produced in accordance with council directive 89/556/EEC								
1. Country of provenance and competent authority	ealth certificate No							
	A. ORIGIN OF EMBRYOS							
3. Approval number of the embryo collection team	3. Approval number of the embryo collection team or embryo production team (¹):							
4. Name and address of the embryo collection production team (¹)	team or embryo	5. Name and address of the consignor						
6. Country and place of loading		7. Means o	of tran	sport				
	B. <b>DESTINATIO</b>	N OF EMBI	RYOS					
8. Member State of destination		9. Name a	nd ado	dress of the consignee				
	C. <b>IDENTIFICATI</b>	ON OF EMI	BRYO	S				
10.1. Identification mark of embryos (²)	10.2. Number of	embryos		10.3. Derived by <i>in vitro</i> fertilisation (a) Subjected to penetration of <i>zona pellucida</i> (b)				
	D. HEALTH I	NFORMATI	ON					
11. I, the undersigned official veterinarian, of the G	overnment of			(name of exporting country)				
certify that:								
11.1. the embryo collection/production team identifie								
<ul> <li>is approved in accordance with Chapter I of Annex A to Directive 89/556/EEC,</li> <li>carried out the collection, processing, or production and storing and transport of the embryos described above in accordance with Chapter II of Annex A to Directive 89/556/EEC,</li> </ul>								
— is subjected at least twice per year to inspection by an official veterinarian.								
11.2. according to official findings								
		(name o	or expo	rting country)				
has: 11.2.1. been free during 12 months immediately prior to collection of the embryos to be exported from rinderpest; 11.2.2. either (¹):								
practise vaccination against it	ng the 12 months in	nmediately p	rior to	collection of the embryos to be exported and does not				
or								

## **▼**M5

- 11.2.2.2. has not been free from foot-and-mouth disease for the 12 months immediately prior to the collection of the embryos and/or practises vaccination against it and
- the donor females and the donors of ovaries, oocytes and other tissues used in the production of embryos come from a holding in which no animal has been vaccinated against foot-and-mouth disease during the 30 days prior to collection, and
- the embryos have been stored in approved conditions for a minimum period of 30 days immediately after collection;

#### 11.2.3. either (1):

has been free from bluetongue and epizootic haemorrhagic disease (EHD) for the 12 months immediately prior to collection of the embryos to be exported and does not practise vaccination against them

or

- 11.2.3.2. has not been free from bluetongue and epizootic haemorrhagic disease (EHD) for the 12 months immediately prior to collection of the embryos to be exported and/or practises vaccination against them and
- the embryos have been stored in approved conditions for a minimum period of 30 days immediately after collection, and
- the donor females and the donors of ovaries, oocytes and other tissues used in the production of embryos were subjected with negative results to an agar gel immuno diffusion test and a serum neutralisation test for epizootic haemorrhagic disease antibodies on a blood sample taken not less than 21 days following collection.

#### 11.3.

- 11.3.1. the premises on which the embryos to be exported or the ovaries, oocytes and other tissues used in the production of embryos to be exported were collected and processed was at the time of collection situated in the centre of an area of 20 km diameter in which according to official findings there had been no incidence of foot-and-mouth disease, bluetongue, epizootic haemorrhagic disease, contagious vesicular stomatitis, Rift Valley fever or contagious bovine pleuropneumonia for 30 days immediately prior to collection and in the case of embryos certified under 11.2.2.2 and 11.2.3.2 for 30 days after collection;
- 11.3.2. between the time of collection or production of the embryos to be exported and their dispatch, they were stored continuously in approved premises which were situated in the centre of an area of 20 km in diameter in which according to official findings there was no incidence of footand-mouth disease, contagious vesicular stomatitis or Rift Valley fever.
- 11.4. the donor females and the donors of ovaries, oocytes and other tissues used in the production of embryos:
- 11.4.1. during the 30 days immediately prior to collection of the embryos to be exported, were located in premises situated in the centre of an area of 20 km in diameter in which according to official findings there was no incidence of foot-and-mouth disease, blue tongue, epizootic haemorrhagic disease, contagious vesicular stomatitis, Rift Valley fever or contagious bovine pleuropneumonia;
- 11.4.2. showed no clinical sign of disease on the day of collection;

11.4.3.	have spent the	e six month	s immediately	prior to	collection	in the	territory	of	
									(name of exporting country)

in a maximum of two herds which are:

- according to official findings free from tuberculosis,
- according to official findings free from brucellosis,
- free from enzootic bovine leukosis or a herd or herds which has/have shown no clinical signs of enzootic bovine leukosis during the previous three years,
- a herd or herds which has/have shown no clinical sign of infectious bovine rhinotracheitis/infectious pustular vulvo-vaginitis during the previous 12 months.
- 11.5. the embryos to be exported were conceived as a result of artificial insemination or in vitro fertilisation with semen complying with the requirements of Council Directive 88/407/EEC and coming from semen collection or storage centres originating in Member States or third countries, in accordance with Article 9(1) of Directive 88/407/EEC, and listed in the Commission's website http://europa.eu.int/comm/food/index\_en.htm.

E. VALIDITY									
12. Date and place	13. Name and qualification of the official veterinarian	14. Signature and stamp of the official veterinarian							
(¹) Delete as necessary. (²) Corresponding to the identification of the donor cows :	and date of collection.								

## PART II

# List of countries approved to use the model animal health certificate at Part I of Annex A

Argentina

Bosnia-Herzegovina

Canada

Croatia

Israel

New Zealand

Romania

Switzerland

United States of America

Former Yugoslav Republic of Macedonia

**▼**<u>M1</u>

**▼**<u>M5</u>

# Part I

ANNEX B

	VETEDINADV	CEDTIEICA	те				
VETERINARY CERTIFICATE embryos of domestic animals of the bovine species for imports collected or produced in accordance with council directive 89/556/EEC							
1. Country of provenance and competent authority	2. H	ealth certificate No					
	A. ORIGIN O	OF EMBRYO	os				
3. Approval number of the embryo collection team of	or embryo producti	ion team (1):					
4. Name and address of the embryo collection production team $(^{\rm I})$	team or embryo	5. Name and address of the consignor					
6. Country and place of loading		7. Means o	of tran	sport			
	B. DESTINATIO	N OF EMBI	RYOS				
8. Member State of destination		9. Name a	nd ado	dress of the consignee			
	C. <b>IDENTIFICATI</b>	ON OF EMI	BRYO	S			
10.1. Identification mark of embryos (²)	10.2. Number of	embryos		10.3. Derived by in vitro fertilisation (a) Subjected to penetration of zona pellucida (b)			
	D. HEALTH II	NFORMATI	ON				
11. I, the undersigned official veterinarian, of the G	overnment of						
				(name of exporting country)			
certify that:							
11.1. the embryo collection/production team identifie							
— is approved in accordance with Chapter I of Ann			41	shower described above in accordance with Chapter II of			
Annex A to Directive 89/556/EEC,	m and storing and	transport of	tne en	nbryos described above in accordance with Chapter II of			
— is subjected at least twice per year to inspection by an official veterinarian.							
11.2. according to official findings							
		(name o	of expo	orting country)			
has:							
11.2.1. been free during 12 months immediately price	or to collection of t	the embryos	to be	exported from rinderpest;			
11.2.2. either (¹):							
11.2.2.1. been free from foot-and-mouth disease during the 12 months immediately prior to collection of the embryos to be exported and does not practise vaccination against it							

## **▼**M5

- 11.2.2.2. has not been free from foot-and-mouth disease for the 12 months immediately prior to the collection of the embryos and/or practises vaccination against it and
- the donor females and the donors of ovaries, oocytes and other tissues used in the production of embryos come from a holding in which no animal has been vaccinated against foot-and-mouth disease during the 30 days prior to collection, and
- the embryos have been stored in approved conditions for a minimum period of 30 days immediately after collection;

#### 11.2.3. either (1):

has been free from bluetongue and epizootic haemorrhagic disease (EHD) for the 12 months immediately prior to collection of the embryos to be exported and does not practise vaccination against them

or

- 11.2.3.2. has not been free from bluetongue and epizootic haemorrhagic disease (EHD) for the 12 months immediately prior to collection of the embryos to be exported and/or practises vaccination against them and
- the embryos have been stored in approved conditions for a minimum period of 30 days immediately after collection, and
- the donor females and the donors of ovaries, oocytes and other tissues used in the production of embryos were subjected with negative results to an agar gel immuno diffusion test and a serum neutralisation test for epizootic haemorrhagic disease antibodies on a blood sample taken not less than 21 days following collection.

#### 11.3.

- 11.3.1. the premises on which the embryos to be exported or the ovaries, oocytes and other tissues used in the production of embryos to be exported were collected and processed was at the time of collection situated in the centre of an area of 20 km diameter in which according to official findings there had been no incidence of foot-and-mouth disease, bluetongue, epizootic haemorrhagic disease, contagious vesicular stomatitis, Rift Valley fever or contagious bovine pleuropneumonia for 30 days immediately prior to collection and in the case of embryos certified under 11.2.2.2 and 11.2.3.2 for 30 days after collection;
- 11.3.2. between the time of collection or production of the embryos to be exported and their dispatch, they were stored continuously in approved premises which were situated in the centre of an area of 20 km in diameter in which according to official findings there was no incidence of footand-mouth disease, contagious vesicular stomatitis or Rift Valley fever.
- 11.4. the donor females and the donors of ovaries, oocytes and other tissues used in the production of embryos:
- 11.4.1. during the 30 days immediately prior to collection of the embryos to be exported, were located in premises situated in the centre of an area of 20 km in diameter in which according to official findings there was no incidence of foot-and-mouth disease, blue tongue, epizootic haemorrhagic disease, contagious vesicular stomatitis, Rift Valley fever or contagious bovine pleuropneumonia;
- 11.4.2. showed no clinical sign of disease on the day of collection;

11.4.3	3. have	spent	the six	months	immediately	prior to	o collection	in t	he	territory	of	
												(name of exporting country)

in a maximum of two herds which are:

- according to official findings free from tuberculosis,
- according to official findings free from brucellosis,
- free from enzootic bovine leukosis or a herd or herds which has/have shown no clinical signs of enzootic bovine leukosis during the previous three years,
- a herd or herds which has/have shown no clinical sign of infectious bovine rhinotracheitis/infectious pustular vulvo-vaginitis during the previous 12 months.
- 11.4.4. were subjected to a serum neutralisation test for Akabane on a blood sample taken not less than 21 days following collection.
- 11.5. the embryos to be exported were conceived as a result of artificial insemination or in vitro fertilisation with semen complying with the requirements of Council Directive 88/407/EEC and coming from semen collection or storage centres originating in Member States or third countries, in accordance with Article 9(1) of Directive 88/407/EEC, and listed in the Commission's website http://europa.eu.int/comm/food/index\_en.htm.

E. VALIDITY								
12. Date and place	13. Name and qualification of the official veterinarian	14. Signature and stamp of the official veterinarian						
(I) Delete as passessory								

- (1) Delete as necessary.
- (2) Corresponding to the identification of the donor cows and date of collection.

# **▼**<u>M1</u>

# PART II

List of countries approved to use the model animal health certificate at Part I of Annex B

Australia