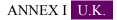
Commission Decision of 7 February 2008 amending Annex D to Council Directive 88/407/EEC and Commission Decision 2004/639/EC laying down the importation conditions of semen of domestic animals of the bovine species (notified under document number C(2008) 409) (Text with EEA relevance) (2008/120/EC)

Changes to legislation: Commission Decision of 7 February 2008 amending Annex D to Council Directive 88/407/EEC and Commission Decision 2004/639/EC laying down the importation conditions of semen of domestic animals of the bovine species (notified under document number C(2008) 409) (Text with EEA relevance) (2008/120/EC) is up to date with all changes known to be in force on or before 02 January 2024. There are changes that may be brought into force at a future date. Changes that have been made appear in the content and are referenced with annotations. (See end of Document for details) View outstanding changes





MODELS OF CERTIFICATES FOR INTRA-COMMUNITY TRADE

ANNEX D1 U.K.

Model of certificate applicable to intra-Community trade in semen collected in accordance with Council Directive 88/407/EEC, as amended by Directive 2003/43/EC, dispatched from an approved semen collection centre

EU	JROPEAN COMMUNITY Intra-Community trade certificate						
Г		Consignor		I.2. Certificate reference number I.2.a. Local reference number			
	l	Name Address		I.3. Central Competent Authority			
Ę	Postal code			I.4. Local Competent Authority			
inte	1.5	Consignee		I.6. No(s) of related original certificates			
rese	ı	Name		No(s) of accompanying documents			
ᇦ		Address					
l iii		Postal code		1.7.			
of consignment presented	1.8.	Country of origin ISO code	I.9. Region of origin Code	I.10. Country of destination ISO code destination Code			
ls o	I.12.	Place of origin		I.13. Place of destination			
Part I: Details		Seme	n centre	Semen centre ☐ Holding ☐			
≓ا		Name	Approval number	Name Approval number			
Par		Address		Address			
		Postal code		Postal code			
	1.14.	Place of loading		I.15. Date of departure			
	Postal code			1.10. Date of departure			
	I.16.	Means of transport		1.17.			
		Aeroplane Ship [Railway wagon				
		Road vehicle	Other				
	1.18.	Description of commodity		I.19. Commodity code (CN code)			
				05 11 10			
				I.20. Number/quantity			
	1.21.	Temperature of products		I.22. Number of packages			
		Ambient	Chilled	Frozen			
	1.23.	Identification of container/seal	number	I.24. Type of packaging			
	1.25.	Commodities certified for: Artificial reproduction]				
	1.26.	Transit through third country		I.27. Transit through Member States			
		Third country	ISO code	Member State ISO code			
		Exit point	Code	Member State ISO code			
		Entry point	BIP unit No	Member State ISO code			
	1.28.			1.29.			
	Third country ISO code						
		Exit point	Code				
	1.30.						
	I.31.	Identification of the commodities					
		Species (Scientific name)	Iden	ification mark Quantity			

EUROPEAN COMMUNITY Bovine semen II.a. Certificate reference number II.b. Local reference number II.1. Animal health attestation I, the undersigned official veterinarian, hereby certify that: II.1.1. The semen described above: (a) was collected, processed and stored under conditions which comply with the standards laid down in Directive 88/407/EEC; Part II: Certification (b) was sent to the place of loading in a sealed container under conditions which comply with Directive 88/407/EEC and bearing the number detailed in Part I.23; II.1.2. The semen described above was collected from bulls, which: (1) either [have not been vaccinated against foot-and-mouth disease within 12 months prior to collection;] [have been vaccinated against foot-and-mouth disease less than 12 months and more than 30 days prior to the collection, and 5 % of doses of semen of each collection, with a minimum of five straws, have been submitted to a virus isolation test for foot-and-mouth disease, carried out with negative results in the laboratory (......) (2) situated in or designated by the Member State of destination:1 The semen described above was stored in approved conditions for a minimum period of 30 days immediately following collection (3). II.1.3. Notes Part I - Box I.12: place of origin shall correspond to the semen collection centre (as defined in Article 2(b) first indent of Directive 88/407/EEC) of semen origin. - Box I.13: place of destination shall correspond to the semen collection or storage centre (as defined in Article 2(b) of Directive 88/407/EEC), or to the holding of semen destination. - Box I.23: identification of container and seal number shall be indicated. - Box I.31: identification mark shall correspond to the identification of the donor animals and the date of collection. (1) Delete as appropriate. (2) Name of the laboratory. (3) May be deleted for fresh semen. - The colour of the stamp and signature must be different from that of the other particulars in the certificate. Official veterinarian or official inspector Name (in capital letters): Qualification and title: Local Veterinary Unit: No of the related LVU: Stamp



Model of certificate applicable from 1 January 2006 to intra-Community trade in stocks of semen collected, processed and/or stored before 31 December 2004 in conformity with the provisions of Council Directive 88/407/EEC, applying

Changes to legislation: Commission Decision of 7 February 2008 amending Annex D to Council Directive 88/407/EEC and Commission Decision 2004/639/EC laying down the importation conditions of semen of domestic animals of the bovine species (notified under document number C(2008) 409) (Text with EEA relevance) (2008/120/EC) is up to date with all changes known to be in force on or before 02 January 2024. There are changes that may be brought into force at a future date. Changes that have been made appear in the content and are referenced with annotations. (See end of Document for details) View outstanding changes

until 1 July 2003 and traded after that date in accordance with Article 2(2) of Directive 2003/43/EC, dispatched from an approved semen collection centre

EUF	ROPEAN COMMUNITY			Intra-Community trade certificate	
	I.1. Consignor		I.2. Certificate reference number	I.2.a. Local reference number	
	Name				
	Address		I.3. Central Competent Authority		
	Postal code		I.4. Local Competent Authority		
ted	I.5. Consignee		I.6. No(s) of related original certification	ntes	
sen	Name		No(s) of accompanying docume		
ore:	Address				
ŧ	Postal code				
consignment presented			1.7.		
ısig	I.8. Country of origin ISO code I.9. Re	egion of origin Code	I.10. Country of ISO code	I.11. Region of Code	
8	iio. Godiniy di Grigin	,	destination	destination	
6					
ails	I.12. Place of origin		I.13. Place of destination		
Part I: Details	Semen centre	. 🗆	Semen centre	Holding	
#	Nome	royal number	Name A	Approval number	
Par		roval number	Address	.,	
	Address				
	Postal code		Postal code		
	I.14. Place of loading		I.15. Date of departure		
	Postal code				
	Lac Manager of Incomment		1.17.		
	I.16. Means of transport		1.17.		
	Aeroplane Ship Ship	Railway wagon			
	Road vehicle Other	Ц			
	I.18. Description of commodity		I.19. Commodity cod	e (CN code)	
				05 11 10	
				I.20. Number/quantity	
	I.21. Temperature of products			I.22. Number of packages	
	Ambient	Chilled	Frozen	1.22. Number of packages	
	_	_	1102011 🗀	104 7 4 4 4	
	I.23. Identification of container/seal number			I.24. Type of packaging	
	I.25. Commodities certified for:				
	Artificial reproduction		T		
	I.26. Transit through third country		I.27. Transit through Member States		
	Third country	ISO code	Member State	ISO code	
	Exit point	Code	Member State	ISO code	
	Entry point	BIP unit No	Member State	ISO code	
	I.28. Export		1.29.		
	Third country ISO code				
	Exit point	Code			
	1.30.				
	I.31. Identification of the commodities				
	Species	Identifica	ation mark	Quantity	
	(Scientific name)				

EUROPEAN COMMUNITY Bovine semen

the content and are referenced with annotations. (See end of Document for details) View outstanding changes

II.a. Certificate reference number

II.1. Animal health attestation

Part II: Certification

- I, the undersigned official veterinarian, hereby certify that:
- II.1.1. The semen described above was collected before the date of 31 December 2004 on a semen collection centre which:
 - (a) was approved under conditions laid down in Chapter I of Annex A to Directive 88/407/EEC;
 - (b) was operated and supervised under conditions laid down in Chapter II of Annex A to Directive 88/407/EEC;
- II.1.2. At the time the semen described above was collected, all bovine animals at the semen collection centre:
 - (a) came from herds and/or were born to dams which satisfy the conditions of points 1 (b) and (c) in Chapter I of Annex B to Directive
 - (b) have, within the 30 days preceding the quarantine isolation period, undergone, with negative results:
 - the tests referred to in points 1(d) (i), (ii) and (iii) of Chapter I of Annex B to Directive 88/407/EEC, and
 - a serum neutralisation test or ELISA test for infectious bovine rhinotracheitis/infectious pustular vulvovaginitis, and
 - a virus isolation test (fluorescent antibody test or immunoperoxidase test) for bovine viral diarrhoea, which in the case of an
 animal less than six months of age has been deferred until that age was reached;
 - (c) have satisfied the quarantine isolation period of 30 days and have been subjected with the required negative results to the following health tests:
 - a serological test for brucellosis carried out in accordance with the procedure described in Annex C to Directive 64/432/EEC;
 - either an immunofluorescent antibody test or a culture test for Campylobacter fetus infection on a sample of preputial material
 or artificial vagina washings, or, in the case of a female animal, a vaginal mucus agglutination test;
 - a microscopic examination and culture test for Trichomonas foetus on a sample of preputial material or artificial vagina washings, or in case of a female animal a vaginal mucus agglutination test;
 - (d) have undergone, at least once a year, with negative results, the routine tests referred to in points 1(a), (b) and (c) in Chapter II of Annex B to Directive 88/407/EEC;
- II.1.3. At the time the semen described above was collected,
 - (a) all female bovine animals in the centre have undergone, at least once a year, a vaginal mucus agglutination test for Campylobacter fetus infection with negative results, and
 - (b) all bulls used for semen production have undergone with negative result either an immunofluorescent antibody test or a culture test for Campylobacter fetus infection on a sample of preputial material or artificial vagina washings carried out within 12 months prior to collection:
- II.1.4. The semen described above was collected from bulls standing in a semen collection centre in which:
- (1) either [all bovine animals have not been vaccinated against infectious bovine rhinotracheitis and have undergone at least once a year with negative result a serum neutralisation test or an ELISA test for infectious bovine rhinotracheitis/infectious pustular vulvovaginitis;]
- (1) or [bovine animals not vaccinated against infectious bovine rhinotracheitis have undergone, at least once a year, with negative result a serum neutralisation test or ELISA test for infectious bovine rhinotracheitis/infectious pustular vulvovaginitis, and testing for infectious bovine rhinotracheitis is not carried out on bulls which have received a first vaccination against infectious bovine rhinotracheitis at the insemination centre after they have been tested with negative result in a serum neutralisation test or ELISA test for infectious bovine rhinotracheitis/infectious pustular vulvovaginitis and which since the first vaccination have been regularly re-vaccinated with an interval of not more than six months;]

Changes to legislation: Commission Decision of 7 February 2008 amending Annex D to Council Directive 88/407/EEC and Commission Decision 2004/639/EC laying down the importation conditions of semen of domestic animals of the bovine species (notified under document number C(2008) 409) (Text with EEA relevance) (2008/120/EC) is up to date with all changes known to be in force on or before 02 January 2024. There are changes that may be brought into force at a future date. Changes that have been made appear in the content and are referenced with annotations. (See end of Document for details) View outstanding changes

EUROPEAN COMMUNITY Bovine semen

II.1.5. The semen described above was collected from bulls which:

II.1.5.1.

- (1) either [have not been vaccinated against foot-and-mouth disease within 12 months prior to collection;]
- [have been vaccinated against foot-and-mouth disease less than 12 months and more than 30 days prior to collection, and 5 % of doses of the semen from each collection, with a minimum of five straws, have been submitted to a virus isolation test for foot-and-mouth disease, carried out with negative results in the laboratory (......) (2), situated in or designated by the Member State of destination;

II.1.5.2.

- (1) either [have not been vaccinated against infectious bovine rhinotracheitis,]
- [have been vaccinated against infectious bovine rhinotracheitis in accordance with point II.1.4,]
- II.1.6. The semen described above was stored in approved conditions for a minimum period of 30 days immediately following collection (3).
- II.1.7. The semen described above was sent to the place of loading in a sealed container and bearing the number detailed in Part I.23.

Notes

Part I

- Box I.12: place of origin shall correspond to the semen collection centre (as defined in Article 2(b) first indent of Directive 88/407/EEC) of semen origin.
- Box I.13: place of destination shall correspond to the semen collection or storage centre (as defined in Article 2(b) of Directive 88/407/EEC), or to the holding of semen destination.
- Box I.23: identification of container and seal number shall be indicated.
- Box I.31: identification mark shall correspond to the identification of the donor animals, the breed of the donor animals, the date of collection which must be earlier than 31 December 2004.

Part II

- (1) Delete as appropriate.
- (2) Name of the laboratory.
- (3) May be deleted for fresh semen.
- The colour of the stamp and signature must be different from that of the other particulars in the certificate.

Official veterinarian or official inspector

Name (in capital letters): Local Veterinary Unit: Date:

Stamp

Qualification and title: No of the related LVU: Signature:

ANNEX Model of certificate applicable to intra-Community trade in semen dispatched from an approved semen storage centre or an approved semen collection centre: D3

- either collected in accordance with Council Directive 88/407/EEC as amended by Directive 2003/43/EC;
- or collected, processed and/or stored before 31 December 2004 in conformity with the provisions of Council Directive 88/407/EEC, applying until 1 July 2003 and traded after that date in accordance with Article 2(2) of Directive 2003/43/EC.

EUF	ROPEAN COMMUNITY			Intra-Community trade certificate	
	I.1. Consignor		I.2. Certificate reference number	I.2.a. Local reference number	
	Name		I.3. Central Competent Authority		
	Address Postal code		1.3. Central Competent Authority		
ted	Postal code		I.4. Local Competent Authority		
sen	I.5. Consignee		I.6. No(s) of related original certification		
t pre	Name Address		No(s) of accompanying docume	nts	
men	Postal code		1.7.		
consignment presented	I.8. Country of origin ISO code	I.9. Region of origin Code	I.10. Country of ISO code destination	I.11. Region of Code destination	
			destination	destination	
Part I: Details of	I.12. Place of origin		I.13. Place of destination		
Deta	Seme	n centre	Semen centre	Holding	
Ξ	Name	Approval number	Name A	pproval number	
Par	Address		Address		
	Postal code		Postal code		
	I.14. Place of loading		I.15. Date of departure		
	Postal code				
	I.16. Means of transport		1.17.		
	Aeroplane 🗌 Ship	☐ Railway wagon ☐			
	Road vehicle	Other			
	I.18. Description of commodity		I.19. Commodity code	e (CN code)	
			0	5 11 10	
				I.20. Number/quantity	
	I.21. Temperature of products			I.22. Number of packages	
	Ambient	Chilled	Frozen		
	I.23. Identification of container/seal	number		I.24. Type of packaging	
	I.25. Commodities certified for:				
	Artificial reproduction [
	I.26. Transit through third country		I.27. Transit through Member States		
	Third country	ISO code	Member State	ISO code	
	Exit point Code		Member State	ISO code	
	Entry point	BIP unit No	Member State	ISO code	
	I.28. Export ISO code Exit point Code		1.29.		
	Exit point	Code			
	1.30.				
	I.31. Identification of the commoditient Species (Scientific name)		ation mark	Quantity	

EUROPEAN COMMUNITY Bovine semen II.a. Certificate reference number II.b. Local reference number II.1. Animal health attestation I, the undersigned official veterinarian, hereby certify that: the semen described above: II.1.1. has been collected, processed and stored for a minimum period of 30 days immediately following collection in an approved semen Part II: Certification collection centre (2) in (1) either [a Member State, operated and supervised in accordance with Chapter I(1) and Chapter II(1) of Annex A to Directive 88/407/EEC;] (1) and/or [a third country listed in Annex I to Decision 2004/639/EC, operated and supervised in accordance with Chapter I(1) and Chapter II(1) of Annex A to Directive 88/407/EEC, and has been imported to the Community under the conditions of Directive 88/407/EEC;] II.1.2. (1) either [was stored in an approved semen storage centre (2) mentioned in Part I.12, operated and supervised in accordance with Chapter I(2) and Chapter II(2) of Annex A to Directive 88/407/EEC;] (1) and/or [was stored in an approved semen collection centre (2) mentioned in Part I.12, operated and supervised in accordance with Chapter I(1) and Chapter II(1) of Annex A to Directive 88/407/EEC:] was sent to the place of loading in a sealed container under conditions which comply with Directive 88/407/EEC and bearing the number detailed in Part I.23. Notes Part I - Box I.6: should correspond to the serial number of the individual official document(s) or health certificate(s) [either INTRA or CVED] that accompanied the semen described above from the approved semen collection centre of its origin to the described above semen storage centre. The original of those documents or those certificates or the officially endorsed copies thereof must be attached to this certificate. - Box I.12: place of origin shall correspond to the semen collection or storage centre (as defined in Article 2(b) of Directive 88/407/EEC) of semen origin. - Box I.13: place of destination shall correspond to the semen collection or storage centre (as defined in Article 2(b) of Directive 88/407/EEC), or to the holding of semen destination. - Box I.23: identification of container and seal number shall be indicated. - Box I.31: identification mark shall correspond to the identification of the donor animals, the breed of the donor animals, the date of collection. (1) Delete as appropriate (2) Only centres listed in accordance with Article 5(2) and 9(1) of Directive 88/407/EEC. http://circa.europa.eu/irc/sanco/vets/info/data/semen/semen.html - The colour of the stamp and signature must be different from that of the other particulars in the certificate. Official veterinarian or official inspector Name (in capital letters): Qualification and title: Local Veterinary Unit: No of the related LVU: Signature: Stamp

ANNEX II U.K.

ANNEX List of third countries from which Member States authorise imports of semen of domestic animals of the bovine speciesOJ L 146, 14.6.1979, p.

15. Annex as last amended by Regulation (EC) No 1791/2006 (OJ L 363, 20.12.2006, p. 1).ISO codeCountryDescription of territory(if appropriate)Additional guaranteesAUAustraliaThe additional guarantees set out in points II.5.4.1.2 and II.5.4.2.2 of the certificate in Part 1 of Annex II are compulsory.CACanadaTerritory as described in Part 1 of Annex I to Council Decision 79/542/EEC.The additional guarantee set out in point II.5.4.1.2 of the certificate in Part 1 of Annex II is compulsory.CHSwitzerlandHRCroatiaNZNew ZealandUSUnited StatesThe additional guarantee set out in point II.5.4.1.2 of the certificate in Part 1 of Annex II is compulsory.

Changes to legislation: Commission Decision of 7 February 2008 amending Annex D to Council Directive 88/407/EEC and Commission Decision 2004/639/EC laying down the importation conditions of semen of domestic animals of the bovine species (notified under document number C(2008) 409) (Text with EEA relevance) (2008/120/EC) is up to date with all changes known to be in force on or before 02 January 2024. There are changes that may be brought into force at a future date. Changes that have been made appear in the content and are referenced with annotations. (See end of Document for details) View outstanding changes

ANNEX II

Model veterinary certificates for imports and transits of semen of domestic animals of the bovine species (for import, collected in accordance with Council Directive 88/407/EEC as amended by Directive 2003/43/EC)

PART 1

Model certificate applicable to imports and transits of semen collected in accordance with Council Directive 88/407/EEC as amended by Directive 2003/43/EC dispatched from an approved semen collection centre

CO	COUNTRY Veterinary certificate to EU					
	1.1.	Consignor	I.2. Certificate reference number I.2.a.			
		Name	I.3. Central Competent Authority			
		Address	1.0. Contra Compount Authority			
l		Tel.	I.4. Local Competent Authority			
len	1.5.	Consignee	I.6. Person responsible for the load in EU			
Į, į		Name	Name			
Suc		Address	Address			
Ö		Postal code	Postal code			
ě		Tel.	Tel.			
of dispatched consignment	1.7.	Country of origin ISO code I.8. Region of origin Code	I.9. Country of ISO code I.10. Region of Code destination			
is o	1.11	1. Place of origin	I.12. Place of destination			
)eta		Name Approval number	Name			
12		Address	Address			
Part I: Details		Name Approval number				
		Address				
		Name Approval number	Postal code			
		Address				
	1.13	3. Place of loading	I.14. Date of departure			
	1.15	5. Means of transport	I.16. Entry BIP in EU			
		Aeroplane ☐ Ship ☐ Railway wagon ☐				
		Road vehicle Other				
	Ide	ntification:	1.17.			
	Do	cumentary references:				
	1.18	3. Description of commodity	I.19. Commodity code (HS code)			
			05 11 10			
			I.20. Quantity			
	1.21	1.	I.22. Number of packages			
	1.23	3. Identification of container/Seal number	1.24.			
	1.25	5. Commodities certified for:				
		Artificial reproduction				
	1.26	6. For transit through EU to third Country	I.27. For import or admission into EU			
		Third country ISO code				
	1.28	3. Identification of the commodities				
		Species Identifica (Scientific name)	ation mark Quantity			

cor	JNTRY			Bovine semer					
			II.a. Certificate reference number						
	II.	Health information							
		I, the undersigned official veterinarian, hereby certify that:							
	II.1.	(name of c	exporting country) (²)						
Part II: Certification		was free from rinderpest and foot-and-mouth disease during the until its date of dispatch and no vaccination against these disease.							
Certif	II.2.	The centre at which the semen to be exported was collected:							
Part II:	II.2.1.	meets the conditions laid down in Chapter I(1) of Annex A $\ensuremath{\text{to}}$	o Council Directive 88/407/EEC;						
	II.2.2.	is operated and supervised in accordance with the conditions	laid down in Chapter II(1) of Annex	A to Council Directive 88/407/EEC.					
	II.3.	The centre at which the semen to be exported was collected was free from rabies, tuberculosis, brucellosis, anthrax and contagious bovine pleuropneumonia during 30 days prior to the date of collection of the semen to be exported and the 30 days after collection (in the case of fresh semen until the day of dispatch).							
	II.4.	The bovine animals standing at the semen collection centre:							
	II.4.1.	come from herds and/or were born to dams which satisfy the conditions of paragraph 1(b) and (c) of Chapter I of Annex B to Directive 88/407/EEC;							
\dashv	II.4.2.	underwent the tests required in accordance with paragraph preceding the quarantine isolation period;	1(d) of Chapter I of Annex B to Direction	ective 88/407/EEC in the 28 days					
	II.4.3.	have satisfied the quarantine isolation period and testing requies 88/407/EEC;	rements laid down in paragraph 1(e) o	of Chapter I of Annex B to Directive					
	11.4.4.	have undergone, at least once a year, the routine tests refer	red to in Chapter II of Annex B to Di	irective 88/407/EEC.					
	II.5.	The semen to be exported was obtained from donor bulls w	hich:						
	II.5.1.	satisfy the conditions laid down in Annex C of Directive 88/4	.07/EEC;						
	II.5.2.	have remained							
	(¹) either	[in the exporting country for at least the last six months prior	r to collection of the semen to be exp	ported;]					
	(¹) or	[in the exporting country for at least 30 days prior to the collection of the semen since entry and they were imported from							
	II.5.3.	fulfil the import conditions for bovine semen laid down in the depending on the status of the country or zone of residence		al Animal Health Code of the OIE,					
	II.5.4.	were resident in the exporting country,							
	II.5.4.1.								
	(1) either	[II.5.4.1.1. which according to official findings is free from e	pizootic haemorrhagic disease (EHD)	:1					
	(¹) or	[II.5.4.1.2. in which according to official findings the exist:	negative on two occasions not more isation test for all above-listed serotyp	than 12 months apart to an agar- es of EHD, carried out in approved					

Changes to legislation: Commission Decision of 7 February 2008 amending Annex D to Council Directive 88/407/EEC and Commission Decision 2004/639/EC laying down the importation conditions of semen of domestic animals of the bovine species (notified under document number C(2008) 409) (Text with EEA relevance) (2008/120/EC) is up to date with all changes known to be in force on or before 02 January 2024. There are changes that may be brought into force at a future date. Changes that have been made appear in the content and are referenced with annotations. (See end of Document for details) View outstanding changes

COUNTRY Bovine semen II.5.4.2. (1) either [5.4.2.1. which according to official findings is free from Akabane disease and Aino disease:] [5.4.2.2. and were tested negative on two occasions not more than 12 months apart to a serum neutralisation test for Akabane virus and Aino virus carried out in approved laboratory on samples of blood taken prior to and not less than 21 days following collection of the semen;] II.6. The semen to be exported was collected after the date on which the centre was approved by the competent national authorities of the The semen to be exported was processed, stored and transported under conditions which satisfy the terms of Directive 88/407/EEC. Notes Part I: - Box reference I.6: Person responsible for the load in EU: this box is to be filled in only if it is a certificate for transit commodity. - Box reference I.12: Place of destination: this box is to be filled in only if it is a certificate for transit commodity. Box reference I.23: identification of container and seal number shall be indicated. Box reference I.26 and I.27: fill in according to whether it is a transit or an import certificate. - Box reference I.28: identification mark shall correspond to the identification of the donor animals and the date of collection. Part II: (1) Delete as necessary. (2) Countries listed in Annex I to Decision 2004/639/EC. (3) Standards for EHD virus diagnostic tests are described in the Bluetongue Chapter of the Manual of Diagnostic Tests and Vaccines for Terrestrial - The signature and the stamp must be in a different colour to that of the printing Official veterinarian Name (in capital letters): Qualification and title:

PART 2

Model certificate applicable from 1 January 2005 to imports and transits of stocks of semen collected, processed and stored before 31 December 2004 in conformity with the provisions of Council Directive 88/407/EEC applying until 1

July 2003, and imported after 31 December 2004 in accordance with Article 2(2) of Directive 2003/43/EC, dispatched from an approved semen collection centre

COI	JNTR	Υ						Veterinary cer	tificate to EU
	l.1.	Consignor			I.2. Certificate	reference	e number	I.2.a.	
		Name			I.3. Central Competent Authority				
	Address								
		I.5. Consignee Name Address Postal code			I.4. Local Con	npetent A	uthority		
nent	I.5.				I.6. Person res	sponsible	for the load	in EU	
ign					Name				
ons					Address				
o p					Postal cod	de			
tche		Tel.		Tel.					
of dispatched consignment	1.7.	Country of origin ISO code	I.8. Region of origin	Code	I.9. Country of destination		ISO code	I.10. Region of destination	Code
Part I: Details of	l.11.	Place of origin			I.12. Place of	destination	on		
Deta		Name	Approval number		Name				
Ξ		Address			Address				
Par		Name	Approval number						
		Address							
		Name Address	Approval number		Postal co	ode			
	I.13.	Place of loading			I.14. Date of departure				
	l.15.	Means of transport			I.16. Entry BIP in EU				
		Aeroplane Ship Ship		I	I.17.				
	lalam	_	Other						
		tification: umentary references:							
					L19 Commodity code (HS code)				
	1.10.	Description of commodity				I.19. Commodity code (HS code) 05 11 10			
					l				
								I.20. Quantity	
	I.21.						I.22. Number of packages		
	I.23. Identification of container/Seal number							1.24.	
	1.25.	Commodities certified for:							
	Artificial reproduction I.26. For transit through EU to third Country ISO code I.27. For i								
					I.27. For impor	rt or adm	ission into E	v 🗖	
	1.28.	Identification of the commodities							
		Species (Scientific name)		Identifica	ation mark			Quantity	

	,,,,,,,,			Boville delliel					
			II.a. Certificate reference number						
	II. Health information								
		I, the undersigned official veterinarian, hereby certify that:							
	II.1.								
		(name of	exporting country) (2)						
Part II: Certification		has been free from rinderpest and foot-and-mouth disease during the 12 months immediately prior to collection of the semen for export and until its date of dispatch and no vaccination against these diseases has taken place during the same period.							
: Cert	II.2.	The semen described above was collected before 31 December 2004 at the semen collection centre which:							
art II	II.2.1.	meets the conditions laid down in Chapter I of Annex A to Directive 88/407/EEC;							
-	II.2.2.	is operated and supervised in accordance with the conditions laid down in Chapter II of Annex A to Directive 88/407/EEC.							
	II.3.	The centre at which the semen to be exported was collected was free from rabies, tuberculosis, brucellosis, anthrax and contagious bovine pleuropneumonia during the period commencing 30 days prior to the date of collection of the semen to be exported and the 30 days after collection.							
	11.4.	At the time semen described above was collected, all bovine animals standing at the semen collection centre:							
	II.4.1.	came from herds and/or were born to dams which satisfy the conditions of paragraph 1(b) and (c) of Chapter I of Annex B to Directive 88/407/EEC;							
\dashv	II.4.2.	had tested negative, within the 30 days preceding the quara	antine isolation period, to:						
		— the tests referred to in points 1(d)(i), (ii) and (iii) of Chap	ter I of Annex B to Directive 88/407/I	EEC, and					
		— a serum neutralisation test or an ELISA test for infectious	s bovine rhinotracheitis/infectious pus	tular vulvo-vaginitis, and					
		 a virus isolation test (fluorescent antibody test or immunoperoxidase test) for bovine viral diarrhoea, deferred until the animal reached the age of six months in the case of younger animals; 							
	II.4.3.	had undergone the 30-day quarantine isolation period and had tested negative to the following health tests:							
		— a serological test for brucellosis carried out in accordance with the procedure described in Annex C to Directive 64/432/EEC;							
		 either an immunofluorescent antibody test or a culture test artificial vagina washings, or, in the case of a female animal. 							
		 a microscopic examination and culture test for <i>Trichomon</i> or in the case of a female animal a vaginal mucus agglu 		aterial or artificial vagina washings,					
	11.4.4.	had tested negative, at least once a year, to the routine tests referred to in points 1(a), (b) and (c) of Chapter II of Annex B to Directive 88/407/EEC.							
	II.5.	At the time the semen described above was collected,							
	II.5.1.	all female bovine animals in the centre had tested negative at least once a year to a vaginal mucus agglutination test for Campylo-bacter fetus infection, and							
	II.5.2.	all bulls used for semen production had tested negative either to an immunofluorescent antibody test or to a culture test for Campylobacter fetus infection on a sample of preputial material or artificial vagina washings carried out in 12 months prior to collection.							
	II.6.	The semen to be exported was obtained from donor bulls w	hich						
	II.6.1.	satisfy the conditions laid down in Annex C of Directive 88/4	407/EEC;						

COUNTRY

Bovine semen

Changes to legislation: Commission Decision of 7 February 2008 amending Annex D to Council Directive 88/407/EEC and Commission Decision 2004/639/EC laying down the importation conditions of semen of domestic animals of the bovine species (notified under document number C(2008) 409) (Text with EEA relevance) (2008/120/EC) is up to date with all changes known to be in force on or before 02 January 2024. There are changes that may be brought into force at a future date. Changes that have been made appear in the content and are referenced with annotations. (See end of Document for details) View outstanding changes

II.6.2. (1) either [were resident in the exporting country during the six months immediately prior to collection of the semen for export;] [were imported from (2) after spending less than six months in the exporting country and at the time of import satisfied the animal health conditions applying to donors of the semen which is intended for export to the Community;] II.6.3. stand in a semen collection centre at which: (1) either [all bovine animals were not vaccinated against infectious bovine rhinotracheitis and tested negative at least once a year to a serum neutralisation test or an ELISA test for infectious bovine rhinotracheitis/infectious pustular vulvo-vaginitis;] (1) or [bovine animals not vaccinated against infectious bovine rhinotracheitis tested negative, at least once a year, to a serum neutralisation test towner animals not vaccinated against infectious bovine rhinotrachelitis/infectious pustular vulvo-vaginitis, at which testing for infectious bovine rhinotrachelitis/infectious pustular vulvo-vaginitis, at which testing for infectious bovine rhinotrachelitis was not carried out on bulls which had received their first vaccination against infectious bovine rhinotrachelitis at the insemination centre after they had tested negative to a serum neutralisation test or an ELISA test for infectious bovine rhinotrachelitis/infectious pustular vulvo-vaginitis and which had been regularly re-vaccinated at intervals of not more than six months since the first vaccination;] II.6.4. (1) either [have not been vaccinated against infectious bovine rhinotracheitis,] (1) or [have been vaccinated against infectious bovine rhinotracheitis in accordance with point II.6.3,] fulfil the import conditions for bovine semen laid down in the Bluetongue Chapter of the Terrestrial Animal Health Code of the OIE, depending on the status of the country or zone of residence;**** II.6.5. II.6.6. taken prior to and not less than 21 days following collection of the semen;** II.6.8. tested negative on two occasions not more than 12 months apart to a serum neutralisation test for Akabane virus carried out in approved laboratory on samples of blood taken prior to and not less than 21 days following collection of the semen. The semen to be exported was collected after the date on which the centre was approved by the competent national authorities of the 11.7. exporting country. II.8. The semen to be exported was processed, stored and transported under conditions which satisfy the terms of Directive 88/407/EEC prior to its amendment by Directive 2003/43/EC. Notes Part I: - Box reference I.6: Person responsible for the load in EU: this box is to be filled in only if it is a certificate for transit commodity. - Box reference I.12: Place of destination: this box is to be filled in only if it is a certificate for transit commodity. Box reference I.23: identification of container and seal number shall be indicated. - Box reference I.26 and I.27: fill in according to whether it is a transit or an import certificate. - Box reference I.28: identification mark shall correspond to the identification of the donor animals and the date of collection that must be prior to

Changes to legislation: Commission Decision of 7 February 2008 amending Annex D to Council Directive 88/407/EEC and Commission Decision 2004/639/EC laying down the importation conditions of semen of domestic animals of the bovine species (notified under document number C(2008) 409) (Text with EEA relevance) (2008/120/EC) is up to date with all changes known to be in force on or before 02 January 2024. There are changes that may be brought into force at a future date. Changes that have been made appear in the content and are referenced with annotations. (See end of Document for details) View outstanding changes

COUNTRY Bovine semen Part II: Delete as necessary Countries listed in Annex I to Decision 2004/639/EC. Standards for EHD virus diagnostic tests are described in the Bluetongue Chapter of the Manual of Diagnostic Tests and Vaccines for **** To be used only by Australia, Canada and the USA. To be used only by Australia and the USA. To be used only by Canada. To be used only by Australia. The signature and the stamp must be in a different colour to that of the printing. Official veterinarian Name (in capital letters): Qualification and title: Signature:

PART 3

Model certificate applicable to imports and transits of semen dispatched from an approved semen storage centre or an approved semen collection centre:

- either collected and processed in accordance with the conditions of Council Directive 88/407/EEC as amended by Directive 2003/43/EC;
- or collected, processed and stored before 31 December 2004 in conformity with the provisions of Council Directive 88/407/EEC applying until 1 July 2003, and imported after 31 December 2004 in accordance with Article 2(2) of Directive 2003/43/EC.

COL	UNTRY Veterinary certificate to EU						
	I.1. Consignor	I.2. Certificate reference number I.2.a.					
	Name Address	I.3. Central Competent Authority					
	Tel.	I.4. Local Competent Authority					
ent	I.5. Consignee	I.6. Person responsible for the consignment in EU					
	Name	Name					
ısig	Address	Address					
8	Postal code	Postal code					
hed	Tel.	Tel.					
Part I: Details of dispatched consignment	I.7. Country of origin ISO code I.8. Region of origin Code	I.9. Country of ISO code I.10. Region of Code destination					
ls of	I.11. Place of origin	I.12. Place of destination					
etai	Name Approval number	Name					
<u>:</u>	Address	Address					
art	Name Approval number						
-	Address						
	Name Approval number	Postal code					
	Address						
	I.13. Place of loading	I.14. Date of departure					
	I.15. Means of transport	I.16. Entry BIP in EU					
	Aeroplane ☐ Ship ☐ Railway wagon ☐						
	Road vehicle Other	I.17. No(s) of related original certificates					
	Identification: Documentary references:						
	·	I.19. Commodity code (HS code)					
	I.18. Description of commodity	05 11 10					
		I.20. Quantity					
	I.21.	I.22. Number of packages					
	I.23. Identification of container/Seal number	1.24.					
I.25. Commodities certified for: Artificial reproduction □							
	I.26. For transit through EU to third Country ISO code	I.27. For import or admission into EU					
	I.28. Identification of the commodities						
	Species Identifica (Scientific name)	ation mark Quantity					

CUL	MIRT			Bovine semei				
			II.a. Certificate reference number					
	II.	Health information	Health information					
		I, the undersigned official veterinarian of		, hereby certify that:				
		(name of	exporting country) (2)					
	II.1.	The centre at which the semen to be exported to the Community was stored:						
ation	(¹) either							
Part II: Certification) of Annex A to Council Directive							
art II:	(¹) or	[II.1.1 meets the conditions laid down in Chapter I(2) of A	nnex A to Directive 88/407/EEC;					
<u>"</u>	and II.1.2 is operated and supervised in accordance with the conditions laid down in Chapter II(2) of Annex A to Co 88/407/EEC.]							
	II.2.	The semen to be exported to the Community:						
II.2.1. has been collected, processed and stored for a minimum period of 30 days immediately following collection in a collection centre (3) operated and supervised in accordance with Chapter I(1) and Chapter II(1) of Annex A to Direction and								
	(1) either	[located in the exporting country;]						
\dashv	(1) and/o	r [located in						
	and	has been imported to the exporting country under condition Community in accordance with Directive 88/407/EEC,]	ns at least as strict as for imports of	semen of bovine species into the				
	II.2.2.	was stored under conditions which satisfy the terms of Dir	ective 88/407/EEC;					
	II.2.3.	was sent to the place of loading in a sealed container und number detailed in Part I.23.	der conditions which comply with Dire	active 88/407/EEC and bearing the				
	Notes							
	Part I:							
	— Вох г	eference I.6: Person responsible for the load in EU: this box	is to be filled in only if it is a certifica	ate for transit commodity.				
	— Вох г	eference I.12: Place of destination: this box is to be filled in	only if it is a certificate for transit con	nmodity.				
	 Box reference I.17: should correspond to the serial number of the individual official document(s) or health certificate(s) that accompanied th semen described above from the approved semen collection centre of its origin to the described above semen storage centre. The original(s of those document(s) or those certificate(s) or the officially endorsed copies of thereof must be attached to this certificate. 							
	— Вох г	eference I.23: identification of container and seal number sha	Il be indicated.					
	— Box r	eference I.26 and I.27: fill in according to whether it is a tran	sit or an import certificate.					
	— Box reference I.28: identification mark shall correspond to the identification of the donor animals and the date of collection.							

CC	OUNTRY					
	Par	t II:				
	(¹)	Delete as necessary.				
	(²)	Countries listed in Annex I to Decision	2004/639/EC and the Member States.			
	(³)	Only centres listed in accordance with A http://circa.europa.eu/irc/sanco/vets/info/o	Article 5(2) and 9(1) of Directive 88/407/EEC data/semen/semen.html			
	_	The signature and the stamp must be in	n a different colour to that of the printing.			
	Offi	cial veterinarian				
		Name (in capital letters):		Qualification and title:		
		Date:	Place:	Signature:'		

Changes to legislation:

Commission Decision of 7 February 2008 amending Annex D to Council Directive 88/407/EEC and Commission Decision 2004/639/EC laying down the importation conditions of semen of domestic animals of the bovine species (notified under document number C(2008) 409) (Text with EEA relevance) (2008/120/EC) is up to date with all changes known to be in force on or before 02 January 2024. There are changes that may be brought into force at a future date. Changes that have been made appear in the content and are referenced with annotations. View outstanding changes

Changes and effects yet to be applied to:

Decision partial repeal by EUDN 2011/630 Decision