Commission Implementing Decision of 18 July 2012 amending Implementing Decision 2011/630/EU as regards animal health requirements relating to bluetongue and Simbu viruses (notified under document C(2012) 4882) (Text with EEA relevance) (2012/415/EU) Changes to legislation: There are currently no known outstanding effects for the Commission Implementing Decision of 18 July 2012 amending Implementing Decision 2011/630/EU as regards animal health requirements relating to bluetongue and Simbu viruses (notified under document C(2012) 4882) (Text with EEA relevance) (2012/415/EU). (See end of Document for details)

ANNEX

1. Annex I is replaced by the following:

- 'ANNEXList of third countries or parts thereof from which Member States shall authorise Ι imports of semen of domestic animals of the bovine speciesThe certificate to be used for imports from Canada is set out in Commission Decision 2005/290/EC of 4 April 2005 on simplified certificates for the importation of bovine semen and fresh pig meat from Canada and amending Decision 2004/639/EC (only for the semen collected in Canada) 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, as approved by Council Decision 1999/201/EC. The certificates to be used for imports from Switzerland are set out in Annex D to Directive 88/407/ EEC, with the adaptations set out in point 4 of Chapter VII(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.'ISO CodeName of the third countryRemarksDescription of the territory(if appropriate)Additional guaranteesAUAustraliaThe additional guarantee concerning testing set out in point II.5.4.1 of the certificate in Section A of Part 1 of Annex II is compulsory.CACanadaCHSwitzerlandCLChileGLGreenlandHRCroatiaISIcelandNZNew ZealandPMSaint Pierre and MiquelonUSUnited StatesThe additional guarantee set out in point II.5.4.1 of the certificate in Section A of Part 1 of Annex II is compulsory.
- 2. In Part 1 of Annex II, Section A is replaced by the following:

SECTION A

Model 1 — Animal health certificate applicable to imports and transits of semen of domestic animals of the bovine species collected, processed and stored in

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accordance with Council Directive 88/407/EEC, as amended by Directive 2003/43/ EC, dispatched from a semen collection centre where the semen was collected

cou	DUNTRY Veterinary certificate to EU								
	l.1.	Consignor Name	I.2. Certificate reference No I.2.a.						
		Address	I.3. Central competent authority						
		Tel.	I.4. Local competent authority						
lent	I.5.	Consignee	I.6. Person responsible for the load in EU Name						
gun		Name							
onsi		Address	Address						
ð		Postal code	Postal code						
tche		Tel.	Tel.						
f 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						
ls of	1 1 1	Place of origin	I.12. Place of destination						
etai	1. 1 1.	-							
Part I : Details		Name Approval number Address	Name Address						
Pa		Name Approval number Address							
		Name Approval number Address	Postal code						
	I.13.	Place of loading	I.14. Date of departure						
	l.15.	Means of transport	I.16. Entry BIP in EU						
		Aeroplane 🗌 Ship 🗌 Railway wagon 🗌							
		Road vehicle Other							
		Identification	1.17.						
		Documentary references							
	l.18.	Description of commodity	I.19. Commodity code (HS code) 05 11 10						
			I.20. Quantity						
	1.21.		I.22. Number of packages						
		Seal/Container No	1.24.						
	I.25. Commodities certified for:								
	1.26.	For transit through EU to third country	I.27. For import or admission into EU						
		Third country ISO code							
	1.28.	Identification of the commodities							
		Species Breed Donor identity ((Scientific name)	Date of collection Approval number of the centre Quantity						

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(COUNTR	1	Bovine semen —					
	II. I	Health information	II.a. Certificate reference No	II.b.				
		I, the undersigned official veterinarian, hereby certify the	at :					
	II.1.	(name of exporting country) (²)						
		was free from rinderpest and foot-and-mouth disease du until its date of dispatch to the Union and no vaccinatio						
5 II.2. The centre (³) described in Box. I.11. at which the semen to be exported was collected:								
Part II: Certification		II.2.1. meets the conditions laid down in Chapter I(1) σ	1. meets the conditions laid down in Chapter I(1) of Annex A to Directive 88/407/EEC;					
ŭ		II.2.2. is operated and supervised in accordance with the conditions laid down in Chapter II(1) of Annex A to Directive 88/4						
Par	II.3.		centre at which the semen to be exported was collected was free from rabies, tuberculosis, brucellosis, anthrax and contagious ne pleuropneumonia during 30 days prior to the date of collection of the semen to be exported and the 30 days after collection (in the of fresh semen until the day of dispatch to the Union).					
	II.4.	The bovine animals standing at the semen collection centre:						
		II.4.1. come from herds which satisfy the conditions of paragraph 1(b) of Chapter I of Annex B to Directive 88/407/EEC;						
		II.4.2. come from herds or were born to dams which comply with the conditions of Chapter I.1(c) of Annex B to Directive 8 were tested at the age of at least 24 months in accordance with Chapter II.1(c) of Annex B to that Directive;						
		II.4.3. underwent the tests required in accordance with preceding the quarantine isolation period;	paragraph 1(d) of Chapter I of Annex B to I	Directive 88/407/EEC in the 28 days				
		II.4.4. have satisfied the quarantine isolation period an Directive 88/407/EEC;	d testing requirements laid down in paragra	aph 1(e) of Chapter I of Annex B to				
		II.4.5. have undergone, at least once a year, the routin	ne tests referred to in Chapter II of Annex E	8 to Directive 88/407/EEC.				
	II.5.	The semen to be exported was obtained from donor bulls which:						
		II.5.1. satisfy the conditions laid down in Annex C of E	Directive 88/407/EEC;					
	(¹) either	[II.5.2. have remained in the exporting country for at le	ast the last six months prior to collection of	the semen to be exported;]				
	(¹) or	[II.5.2. have remained in the exporting country for at i imported from	ing the period of less than six months prior	to the collection of the semen and				
	(¹) either	[II.5.3. were kept in a bluetongue virus-free country or	zone for at least 60 days prior to, and durir	ng, collection of the semen;]				
	(¹) or	[II.5.3. were kept during a bluetongue virus seasonally collection of the semen;]	free period in a seasonally free zone for a	t least 60 days prior to, and during				
	(1) or	[II.5.3. were kept in a vector-protected establishment for	or at least 60 days prior to, and during colle	ction of the semen;]				
	(¹) or	[II.5.3. were subjected to a serological test to detect ar Manual of Diagnostic Tests and Vaccines for Ti collection period and between 21 and 60 days an	errestrial Animals, with negative results, at	least every 60 days throughout the				
	(¹) or	[II.5.3. were subjected to an agent identification test for Tests and Vaccines for Terrestrial Animals with n this consignment of semen and at least every 7 d this consignment of semen;]	egative results on blood samples taken at co	mmencement and final collection for				
		II.5.4. were resident in the exporting country,						
	(†)	either [II.5.4.1. which according to official findings is fi	ree from epizootic haemorrhagic disease (E	HD);]				

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COUNTRY Bovine semen — Section						
II. Health information	II.a. Certificate reference No	II.b.				
	g to official findings the following serotypes of epizo and were subjected with negative results in eac					
	t more than 12 months apart a serological test (4) carried out i to and not less than 21 days following collection for this con					
more than 60 days	[a serological test (⁴) for the detection of antibody to the EHDV group, carried out on samples taken at intervals of no more than 60 days throughout the collection period and between 21 and 60 days after the final collection for thi consignment of semen.]]					
	on test (⁴) carried out in approved laboratories on blood samp t least every 7 days (virus isolation test) or at least every 28 semen.]]					
II.6. The semen to be exported was colle exporting country;	cted after the date on which the centre was approved by the	competent national authorities of the				
II.7. The semen to be exported was proc	essed, stored and transported under conditions which satisfy	y the terms of Directive 88/407/EEC.				
Notes						
Part I:						
Box I.6: Person responsible for the load in E	Box 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.					
Commission website:	ne semen collection centre listed in accordance with Article n_ova/bovine/index_en.htm and where the semen was collect					
Box I.22: number of packages shall correspon	d to the number of containers.					
Box I.23: identification of container and seal n	umber shall be indicated.					
Box I.26: fill in according to whether it is a tra	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.						
Donor identity shall correspond to the Date of collection shall be indicated		entre indicated in Box I.11 where the				
Part II:						
(1) Delete as necessary.						
(²) Only third countries listed in Annex I to Implementing Decision 2011/630/EU.						
(³) Only semen collection centres listed in accordance with Article 9(2) of Directive 88/407/EEC on the Commission website: http://ec.europa.eu/food/animal/semen_ova/bovine/index_en.htm						
(4) Standards for EHD virus diagnostic tests are described in the Bluetongue Chapter of the Manual of Diagnostic Tests and Vaccines for Terrestrial Animals.						
(⁵) Compulsory for Australia, Canada and the United States.						
- The signature and the stamp must be in a different colour to that of the printing.						

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	Bovine semen — Section A	
II.a. Certificate reference No	II.b.	
Official veterinarian		
Qualifica	Qualification and title:	
Signatu	Signature:	
	Qualific	

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