Status: Point in time view as at 01/07/2016. Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

Commission Regulation (EU) No 206/2010 of 12 March 2010 laying down lists of third countries, territories or parts thereof authorised for the introduction into the European Union of certain animals and fresh meat and the veterinary certification requirements (Text with EEA relevance)

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

[X1ANNEX I

UNGULATES

Editorial Information

X1 Substituted by Corrigendum to Commission Regulation (EU) No 206/2010 of 12 March 2010 laying down lists of third countries, territories or parts thereof authorised for the introduction into the European Union of certain animals and fresh meat and the veterinary certification requirements (Official Journal of the European Union L 73 of 20 March 2010).

$\label{eq:FIPART 1} \textbf{LIST OF THIRD COUNTRIES, TERRITORIES OR PARTS THEREOF}^0$

ISO code Code o		Description	Veterinary ce	Specific	
and name of third country	Territory	of third country, territory or part thereof	Model(s)	SG	conditions
1	2	3	4	5	6
[^{F2} BD — Bangladesh ^f	BD-0	The area covered by Chittagong Safari Park	TRE-A ^g]
CA – Canada	CA-0	Whole country	POR-X		IVb IX
	CA-1	Whole country, except the Okanagan Valley region of British Columbia described as follows:	BOV-X, OVI-X, OVI- Y RUM ^b	A	V

- **a** Without prejudice to specific certification requirements provided for by any relevant agreement between the Union and third countries.
- **b** Exclusively for live animals other than animals belonging to the cervidae species.
- c Certificates in accordance with the Agreement between the European Community and the Swiss Confederation on trade in agricultural products (OJ L 114, 30.4.2002, p. 132).
- d The former Yugoslav Republic of Macedonia: the definitive nomenclature for this country will be agreed following current negotiations at UN level.
- e Not including Kosovo under UNSCR 1244/99.
- **f** [F2This entry applies until 17 August 2015.
- **g** Exclusively for live ungulates of the *Elephas* ssp. from an approved body, institute or centre in Bangladesh to an approved body, institute or centre in Cyprus.]

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

	From
	a
	point
	on
	the
	Canada/
	United
	States
	border
	120°15′
	longitude,
	49°
	latitude
_	Northerly
	to a
	point
	119°35′
	longitude,
	50°30′
	latitude
	North-
	easterly
	to a
	point
	119°
	longitude,
	50°45′
	latitude
	Southerly
	to a
	point
	on the
	Canada/
	United
	States
	border
	118°15′
	longitude,
	- 0,

- Without prejudice to specific certification requirements provided for by any relevant agreement between the Union and
- b Exclusively for live animals other than animals belonging to the cervidae species.
- Certificates in accordance with the Agreement between the European Community and the Swiss Confederation on trade in agricultural products (OJ L 114, 30.4.2002, p. 132).
- The former Yugoslav Republic of Macedonia: the definitive nomenclature for this country will be agreed following current negotiations at UN level.
- Not including Kosovo under UNSCR 1244/99.
- f [F2This entry applies until 17 August 2015.
- Exclusively for live ungulates of the *Elephas* ssp. from an approved body, institute or centre in Bangladesh to an approved body, institute or centre in Cyprus.]

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

		49° latitu	ıde		
CH – Switzerland	CH-0	Whole country	С		
CL – Chile	CL-0	Whole country	BOV-X,OVI- X, RUM		
			POR-X, SUI	В	
GL – Greenland	GL-0	Whole country	OVI-X, RUM		V
[F3]		•	•		
ISCIceland	IS-0	Whole country	BOV-X, BOV-Y RUM, OVI- X, OVI-Y		
			POR-X, POR-Y	В	
ME – Montenegro	ME-0	Whole country			I
MK – The former Yugoslav Republic of Macedonia ^d	MK-0	Whole country			I
[F4NZ – New Zealand	NZ-0	Whole country	BOV-X, BOV-Y, RUM, POR-X, POR-Y OVI-X, OVI-		III V XII]
PM – St Pierre and Miquelon	PM-0	Whole country	BOV-X, BOV-Y, RUM, OVI-		

- a Without prejudice to specific certification requirements provided for by any relevant agreement between the Union and third countries.
- **b** Exclusively for live animals other than animals belonging to the cervidae species.
- c Certificates in accordance with the Agreement between the European Community and the Swiss Confederation on trade in agricultural products (OJ L 114, 30.4.2002, p. 132).
- d The former Yugoslav Republic of Macedonia: the definitive nomenclature for this country will be agreed following current negotiations at UN level.
- e Not including Kosovo under UNSCR 1244/99.
- **f** [F2This entry applies until 17 August 2015.
- g Exclusively for live ungulates of the *Elephas* ssp. from an approved body, institute or centre in Bangladesh to an approved body, institute or centre in Cyprus.]

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

			X, OVI-Y CAM		
RS – Serbia ^e	RS-0	Whole country			I
RU – Russia	RU-0	Whole country			
	RU-1	Whole country except the region of Kaliningrad			
	RU-2	Region of Kaliningrad	BOV-X- TRANSIT- RU		X
[F5US – United States	US-0	Whole country	POR-X	D	1

- Without prejudice to specific certification requirements provided for by any relevant agreement between the Union and
- Exclusively for live animals other than animals belonging to the cervidae species.
- Certificates in accordance with the Agreement between the European Community and the Swiss Confederation on trade in agricultural products (OJ L 114, 30.4.2002, p. 132).
- d The former Yugoslav Republic of Macedonia: the definitive nomenclature for this country will be agreed following current negotiations at UN level.
- Not including Kosovo under UNSCR 1244/99. e
- f [F2This entry applies until 17 August 2015.
- Exclusively for live ungulates of the Elephas ssp. from an approved body, institute or centre in Bangladesh to an approved body, institute or centre in Cyprus.]

Textual Amendments

- Inserted by Commission Implementing Regulation (EU) 2015/917 of 15 June 2015 amending Annex I to Regulation (EU) No 206/2010 laying down lists of third countries, territories or parts thereof authorised for the introduction into the European Union of certain animals and fresh meat and the veterinary certification requirements as regards Bangladesh (Text with EEA relevance).
- F3 Deleted by Commission Regulation (EU) No 519/2013 of 21 February 2013 adapting certain regulations and decisions in the fields of free movement of goods, freedom of movement for persons, right of establishment and freedom to provide services, company law, competition policy, agriculture, food safety, veterinary and phytosanitary policy, fisheries, transport policy, energy, taxation, statistics, social policy and employment, environment, customs union, external relations, and foreign, security and defence policy, by reason of the accession of Croatia.
- F4 Substituted by Commission Implementing Regulation (EU) 2015/604 of 16 April 2015 amending Annexes I and II to Regulation (EU) No 206/2010 as regards animal health requirements for bovine tuberculosis in the models of veterinary certificates BOV-X and BOV-Y and the entries for Israel, New Zealand and Paraguay in the lists of third countries, territories or parts thereof from which the introduction into the Union of live animals and fresh meat is authorised (Text with EEA relevance).
- F5 Inserted by Commission Implementing Regulation (EU) No 102/2013 of 4 February 2013 amending Regulation (EU) No 206/2010 as regards the entry for the United States in the list of third countries, territories or parts thereof authorised for the introduction of live ungulates into the Union, the model

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

veterinary certificate 'POR-X' and the protocols for testing for vesicular stomatitis (Text with EEA relevance).

Specific Conditions (see footnotes in each certificate)

ʻI'

for transit through the territory of a third country of live animals for immediate slaughter or live bovine animals for fattening which are consigned from a Member State and destined to another Member State in lorries which have been sealed with a serially numbered seal.

The seal number must be entered on the health certificate issued in accordance with the model laid down in Annex F to Directive 64/432/ EEC⁽¹⁾ for live bovine animals for slaughter and fattening and in accordance with Model I of Annex E to Directive 91/68/EEC⁽²⁾ for ovine and caprine animals for slaughter.

In addition, the seal must be intact on arrival at the designated border inspection post of entry into the Union and the seal number recorded in the integrated computerised veterinary system of the Union (TRACES).

The certificate must be stamped at the exit point of the Union by the competent veterinary authority prior to transiting one or more third countries, with the following wording 'ONLY FOR TRANSIT BETWEEN DIFFERENT PARTS OF THE EUROPEAN UNION VIA THE FORMER YUGOSLAV REPUBLIC OF MACEDONIA/MONTENEGRO/SERBIA⁽³⁾⁽⁴⁾'.

Bovine animals for fattening must be transported directly to the holding of destination designated by the competent veterinary authority of destination. Those animals must not be moved from that holding unless for immediate slaughter.

'II'

territory recognised as having an official tuberculosis-free status for the purposes of exports to the Union of live animals certified according to the model of certificate BOV-X.

'III'

: territory recognised as having an official brucellosis-free status for the purposes of exports to the Union of live animals certified according to the model of certificate BOV-X.

'IVa'

territory recognised as having an official enzootic-bovine-leukosis (EBL) free status for the purposes of exports to the Union of live animals certified according to the model of certificate BOV –X.

'IVb'

recognised as having officially enzootic-bovine-leukosis (EBL)-free herds equivalent to the requirements set out in Annex D to Directive 64/432/EEC for the purposes of exports to the Union of live animals certified according to the model of certificate BOV–X.

'V

territory recognised as having an official brucellosis-free status for the purposes of exports to the Union of live animals certified according to the model of certificate OVI-X.

'VI'

Geographical constraints:

'VII'

territory recognised as having an official tuberculosis-free status for the purposes of exports to the Union of live animals certified according to the model of certificate RUM.

'VIII'

territory recognised as having an official brucellosis-free status for the purposes of exports to the Union of live animals certified according to the model of certificate RUM.

ANNEX I PART 2 Document Generated: 2024-06-03

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

'IX' : territory recognised as having an official Aujeszky's disease -free status for the purposes of exports to the Union of live animals certified according to the model of certificate POR-X.

'X' : Only for transit through Lithuania of bovine animals for breeding and/

or production from the Kaliningrad region to other regions of Russia.

'[F6XI']: holdings or compartments recognised as applying controlled housing conditions in accordance with Article 8 of Regulation (EC) No

2075/2005.]

'I^{F7}XII' : territory recognised as having officially tuberculosis-free bovine herds

equivalent to those recognised based on the conditions laid down in paragraphs 1 and 2 of Annex A.I to Directive 64/432/EEC, for the purposes of exports to the Union of live animals certified according to

the model of veterinary certificate BOV-X or BOV-Y.]

Textual Amendments

F6 Inserted by Commission Implementing Regulation (EU) No 1218/2014 of 13 November 2014 amending Annexes I and II to Regulation (EU) No 206/2010 as regards animal health requirements for Trichinella in the model of veterinary certificate for imports into the Union of domestic porcine animals intended for breeding, production or slaughter, and of fresh meat thereof (Text with EEA relevance).

F7 Inserted by Commission Implementing Regulation (EU) 2015/604 of 16 April 2015 amending Annexes I and II to Regulation (EU) No 206/2010 as regards animal health requirements for bovine tuberculosis in the models of veterinary certificates BOV-X and BOV-Y and the entries for Israel, New Zealand and Paraguay in the lists of third countries, territories or parts thereof from which the introduction into the Union of live animals and fresh meat is authorised (Text with EEA relevance).

Textual Amendments

F1 Substituted by Commission Implementing Regulation (EU) No 644/2012 of 16 July 2012 amending Regulation (EU) No 206/2010 laying down lists of third countries, territories or parts thereof authorised for the introduction into the European Union of certain animals and fresh meat and the veterinary certification requirements, as regards Russia (Text with EEA relevance).

PART 2

Models of Veterinary Certificates

	Models of veterinary Certificates
Models	·
'BOV-X'	: Model of veterinary certificate for domestic bovine animals (including Bubalus and Bison species and their cross-breeds) intended for breeding and/or production after importation.
'BOV-Y'	: Model of veterinary certificate for domestic bovine animals (including Bubalus and Bison species and their cross-breeds) intended for immediate slaughter after importation.
'BOV-X-	: Model of veterinary certificate for domestic bovine animals (including
TRANSIT-RU'	Bubalus and Bison species and their cross-breeds) intended for transit from the region of Kaliningrad to other regions of Russia via the territory of Lithuania.
'OVI-X'	: Model of veterinary certificate for domestic ovine animals (Ovis aries) and domestic caprine animals (Capra hircus) intended for breeding and/

or production after importation.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

'OVI-Y' : Model of veterinary certificate for domestic ovine animals (Ovis aries)

and domestic caprine animals (Capra hircus) intended for immediate

slaughter after importation.

'IF8POR-X' : Model of veterinary certificate for domestic porcine animals (Sus

scrofa) intended for breeding and/or production after importation or intended for transit through the Union from one third country to another

third country.]

'POR-Y' : Model of veterinary certificate for domestic porcine animals (Sus

scrofa) intended for immediate slaughter after importation.

'RUM' : Model of veterinary certificate for animals of the order Artiodactyla

(excluding bovine animals (including Bubalus and Bison species and their cross-breeds), Ovis aries, Capra hircus, Suidae and Tayassuidae),

and of the families Rhinocerotidae and Elephantidae.

'SUI' : Model of veterinary certificate for non-domestic Suidae, Tayassuidae

and Tapiridae.

'CAM' : Model of specific attestation for animals imported from St Pierre and

Miquelon under the conditions provided for in Part 7 of Annex I.

Textual Amendments

Substituted by Commission Implementing Regulation (EU) No 102/2013 of 4 February 2013 amending Regulation (EU) No 206/2010 as regards the entry for the United States in the list of third countries, territories or parts thereof authorised for the introduction of live ungulates into the Union, the model veterinary certificate 'POR-X' and the protocols for testing for vesicular stomatitis (Text with EEA relevance).

SG (Supplementary guarantees)

'A' : guarantees regarding Bluetongue and Epizootic-haemorrhagic-disease

tests on animals certified according to the model of veterinary certificates BOV-X (point II.2.8 B), OVI-X (point II.2.6 D) and RUM

(point II.2.6).

'B' : guarantees regarding Swine-vesicular-disease and Classical-swine-

fever tests on animals certified according to the model of veterinary

certificates POR-X (point II.2.4 B) and SUI (point II.2.4 B).

'C' : guarantees regarding Brucellosis test on animals certified according to

the model of veterinary certificates POR-X (point II.2.4 C) and SUI

(point II.2.4 C).

'[F8D' : guarantees regarding vesicular stomatitis test on animals certified

according to the model of veterinary certificate POR-X (point

II.2.1(b)).]]

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

Model BOV-X

COUN	TRY:					Veterinary cer	tificate to El
	1.1.	Consignor	1.2.	Certificate referen	ice No	1.2.a.	
	Name Address		1.3.	Central competen	t authority		
		Tel.	1.4.	Local competent a	authority		
nent	1.5.	Consignee	1.6.				
ignn		Name					
cons		Address					
tched		Postal code Tel.					
Part I: Details of dispatched consignment	1.7.	Country ISO of origin code origin	1.9.	Country of destination	ISO code	I.10. Region of destination	Code
Detail	1.11.	Place of origin	1.12	<u>. </u>			
i.		Name Approval number Address					
ď		Address	_				
	I.13.	Place of loading Address Approval number	1.14	. Date of departure			
	I.15.	Means of transport	1.16	i. Entry BIP in EU			
		Aeroplane 🔲 🚆 Ship 🗖					
		Railway wagon ☐ Road vehicle ☐ Other ☐	1.17				
		Identification					
	118	Documentary references Description of commodity			I 19 Commo	odity code (HS code	\
	1.10.	becomplied of commonly			01.	-	,
						I.20. Quantity	
	1.21.					I.22. Number of p	oackages
		Seal/Container No				1.24.	
	1.25.	Commodities certified for:					
		Breeding		Fattening	, 🗆		
	1.26.			I.27. For import or	admission into	EU 🗖	
	1.28.	Identification of the commodities					
	l .	Species Breed Identi entific name)	ficatio	n system Ider	ntification numb	per Age	Sex

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model BOV-X

	II. I	Health information	on		II.a.	Certificate refer	ence number	II.b.		
	II.1.	Public Health Attestation								
		I, the undersigned official veterinarian, hereby certify, that the animals described in this certificate:								
tion		II.1.1.	past 42 6 mont	2 days in the case	of bru abies,	cellosis, for the p	past 30 days in the cas	n on health grounds, for the e of anthrax and for the past mals from holdings which did		
tifica		II.1.2.	have not received:							
.ce			_	 any stilbene or thyrostatic substances, 						
Part II: Certification			-				β- agonist substance s defined in Directive 9	es for purposes other than 96/22/EC);		
		II.1.3.	with re	gard to bovine spo	ongifor	m encephalopath	ny (BSE):			
		(¹) (²) either	[(a)	back to the dam	and h	erd of origin, an		n enabling them to be traced vine animals as described in) No 999/2001;		
			(b)	after the date fro and greaves der	om whi	ich the ban on t om ruminants had	he feeding of ruminan	erned, the animals were born ts with meat-and-bone meal rced or after the date of birth ed ban.]		
		(¹) (³) or	[(a)	the animals are identified by a permanent identification system enabling them to be trace- back to the dam and herd of origin, and are not exposed bovine animals as described in Chapter C, Part II, point (4)(b)(iv) of Annex II to Regulation (EC) No 999/2001;						
			(b)	b) the animals were born after the date from which the ban on the feeding of ruminants with meat-and-bone meal and greaves derived from ruminants had been effectively enforced or after the date of birth of the last BSE indigenous case if born after the date of the feed ban.]						
		(¹) (⁴) or	[(a)	back to the dam	and h	erd of origin, an		n enabling them to be traced vine animals as described in C) No 999/2001;		
			(b)	ruminants with	meat-a ed or	and-bone meal after the date of	and greaves derived	ch the ban on the feeding of from ruminants had been indigenous case if born after		
	II.2.	2. Animal Health attestation:								
		I, the undersigned requirements:	gned of	ficial veterinarian,	herek	by certify, that t	the animals described	d above meet the following		
		II.2.1.		ome from the territ	ory wit	h code:	(⁵) v	which, at the date of issuing		
		(1) either	[(a)	has been free fo	24 m	onths from foot-a	nd-mouth disease]			
		(¹) or	[(a)	without having	had c	ases/outbreaks	after that date, and	authorised to export these , of (dd/mm/yyyy),]		
			(b)		a, lum	py skin disease		y fever, contagious bovine norrhagic disease, and for		
			(c)		o) has	been carried ou	it and imports of dom	the diseases mentioned in estic cloven-hoofed animals		
		(1) either	[(d)	has been free fo	24 m	onths from blueto	ongue;]			

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model BOV-X

COUN	IKI			Model BOV-X				
II.	Health informati	ion	II.a. Certificate reference number	II.b.				
	(1) (2) or [(d) has been free for 24 months from bluetongue, and the animals have reacted negatively to a serological test for the detection of antibody for bluetongue and epizootic haemorrhagid disease, carried out on two occasions on samples of blood taken at the beginning of the isolation/quarantine period and at least 28 days later, on							
	(¹) or	with an ina against all source popu a 150 km ra	with an inactivated vaccine, at least 60 days before the date of dispatch to the Union, against all bluetongue serotype/s (insert serotype/s) which are those present in the source population as demonstrated through a surveillance programme (12) in an area with a 150 km radius around the holding(s) of origin described under box reference I.11, and the animals are still within the immunity period of time guaranteed in the specifications of					
	II.2.2.		n the territory described under point II.2.1 sind atch to the Union and without contact with im					
	II.2.3.	they have remained described under box	since birth or at least 40 days before dispate	ch in the holding(s) of origin				
			l which, in an area with a 150 km radius, there morrhagic disease during the previous 60 days,					
		foot-and-mo	I which, in an area with a 10 km radius, there th disease, rinderpest, Rift valley fever, blo onia, lumpy skin disease and, vesicular st	uetongue, contagious bovine				
	II.2.4.		to be killed under a national programme for th nated against the diseases referred to under po					
	II.2.5.		ds that are not restricted under the national losis, brucellosis and enzootic bovine leukosis;					
	II.2.6.	they come from her	recognised as officially tuberculosis-free (⁶) (^{6b});				
and	(¹) (⁷) either	[come from a region	which is recognised as officially tuberculosis-free	e (⁶);]				
	(¹) or		I to an intradermal tuberculin test (8) carried or re dispatch to the Union;]	ut with negative results within				
	(¹) or	[are less than 6 wee	s old;]					
	II.2.7.	they have not been brucellosis-free (6);	vaccinated against brucellosis and come from	herds recognised as officially				
and	(1) (7) either	[come from a region	which is recognised as officially brucellosis-free	(⁶),]				
	(¹) or		I to at least one test for bovine brucellosis (8) is before dispatch to the Union,]	carried out on samples taken				
	(¹) or	[are less than 12 mo	ths old,]					
	(¹) or	[are castrated male:	of any age,]					
(¹) ei	ther [II.2.8.	•	included in an official system for the control of en no evidence either clinical or as a result of a 's,]					
(¹) or	[II.2.8.	they come from her	recognised as officially enzootic-bovine-leukos	sis-free (⁶) (^{6a}),]				
and	(¹) (⁷) either	[come from a region	which is recognised as officially enzootic-bovine	-leukosis-free (⁶);]				
	(¹) or		to an individual test for enzootic bovine leukos en within the past 30 days before dispatch to th					
	(¹) or	[are less than 12 mo	ths old;]					
	II.2.9.	they are/were (1) dis	atched from their holding(s) of origin, without pa	assing through any market:				

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model BOV-X

II.	Health information		II.a.	Certificate reference number	II.b.			
	(1) either	[directly	y to the Union,]					
	(¹) or		[to the officially authorised assembly centre described under box reference I.13 situated within the territory described under point II.2.1,]					
		and, ur	ntil dispatched to th	ne Uni	on:			
		(a)	(a) they did not come in contact with other cloven-hoofed animals not complying with the health requirements as described in this certificate,					
		(b)	they were not at any place where, or around which, within a 10 km radius, during the previous 30 days there has been a case/outbreak of any of the diseases referred to in point II.2.1;					
	II.2.10.		•		tainers in which they were loaded we authorised disinfectant;	ere cleaned and disinfected		
	II.2.11.		ere examined by disease;	an offi	cial veterinarian within 24 hours of loa	ding and showed no clinical		
	II.2.12.	they have been loaded for dispatch to the Union on						
II.3.	Animal transport attestation							

I, the undersigned official veterinarian, hereby certify, that the animals described above have been treated before and at the time of loading in accordance with the relevant provisions of Regulation (EC) No 1/2005, in particular as regards watering and feeding, and they are fit for the intended transport.

(1) (11) [II.4. Specific requirements

- According to official information, no clinical or pathological evidence of infectious bovine II.4.1. rhinotracheitis (IBR) has been recorded in the holding(s) of origin referred to in box reference I.11, for the last 12 months;
- 11.4.2. the animals referred to in box reference I.28:
 - have been isolated in accommodation approved by the competent authority for the last 30 days immediately prior to dispatch for export,
 - (b) have been subjected to a serological test for IBR on sera taken at least 21 days after entry into isolation, with negative results, and all animals in isolation have also given negative results to this test.
 - have not been vaccinated against IBR.] (c)

Notes

This certificate is meant for domestic bovine animals (including Bubalus and Bison species and their cross-breeds) intended for breeding and/or production.

After importation the animals must be conveyed without delay to the holding of destination where they shall remain for a minimum period of 30 days before further movement outside the holding, except in the case of a dispatch to a slaughterhouse.

Part I:

_	Box reference I.8:	Provide the code of territory as appearing in Part 1 of Annex I to Regulation (EU) No 206/2010.
-	Box reference I.13:	The assembly centre, if any, must fulfil the conditions for its approval, as laid down in Part 5 of Annex I to Regulation (EU) No $206/2010$.
_	Box reference I.15:	Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor must inform the BIP of entry into the Union.

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model BOV-X

II.	Health information		II.a.	Certificate reference number	II.b.
_	Box reference I.23:	For containers of be included.	boxe	s, the container number and the seal i	number (if applicable) should
_	Box reference I.28:	Identification sys	tem: T	he animals must bear:	
				which permits tracing of their prenuch as tag, tattoos, brand, chip, transpo	0 , ,
		•		les the ISO code of the exporting cou their premises of origin.	untry. The individual number
		Species: Select a	mong	st "Bos", "Bison" and "Bubalus" as app	ropriate.
		Age: Date of birth	ı (dd/n	mm/yyyy).	
		Sex (M = male, F	= fem	nale, C = castrated).	
		Breed: select pur	ebred	, crossbreed.	

Part II:

- (1) Keep as appropriate.
- (2) Only if the animals were born and continuously reared in a country or region categorised in accordance with Article 5(2) of Regulation (EC) No 999/2001 as a country or region posing a negligible BSE risk and listed as such in Decision 2007/453/EC.
- (3) Only if the country or region of origin is categorised in accordance with Article 5(2) of Regulation (EC) No 999/2001 as a country or region posing a controlled BSE risk and is listed as such in Decision 2007/453/EC.
- (d) Only if the country or region of origin has not been categorised in accordance with Article 5(2) of Regulation (EC) No 999/2001 or has been categorised as a country or region with undetermined BSE risk and listed as such in Decision 2007/453/EC.
- (5) Code of the territory as it appears in Part 1 of Annex I to Regulation (EU) No 206/2010
- (6) Officially tuberculosis/brucellosis-free regions and herds as laid down in Annex A to Directive 64/432/EEC; and enzootic-bovine-leukosis-free regions and herds as laid down in Chapter I of Annex D to Directive 64/432/EEC.
- (⁵⁸) Only for officially enzootic-bovine-leukosis-free herds recognised as equivalent to the requirements as laid down in Chapter I of Annex D to Directive 64/432/EEC for the purpose of exports to the EU of live animals according to the model of veterinary certificate BOV-X from the territory that, in column 6 of Part 1 of Annex I to Regulation (EU) No 206/2010, appears with the entry "IVb" as regards enzootic bovine leukosis.
- (^{6b}) Only for a territory appearing with entry "XII" in column 6 of Part 1 of Annex I to Regulation (EU) No 206/2010 indicating that bovine herds officially declared tuberculosis-free are recognised based on equivalent conditions to those laid down in paragraphs 1 and 2 of Annex A.I to Directive 64/432/EEC for the purposes of exports to the Union of live animals certified according to the model of veterinary certificate BOV-X.
- (⁷) Only for a territory that, in column 6 of Part 1 of Annex I to Regulation (EU) No 206/2010, appears with the entry "II", as regards tuberculosis, "III", as regards brucellosis, and/or "IVa" as regards enzootic bovine leukosis.
- (8) Tests carried out in accordance with the protocols that, for the disease concerned, are described in Part 6 of Annex I to Regulation (EU) No 206/2010.
- (°) Supplementary guarantees to be provided when required in column 5 "SG" of Part 1 of Annex I to Regulation (EU) No 206/2010, with the entry "A".
 - Tests for bluetongue and for epizootic haemorrhagic disease in accordance with Part 6 of Annex I to Regulation (EU) No 206/2010.
- (10) Date of loading. Imports of these animals shall not be allowed when the animals were loaded either prior to the date of authorisation for exportation to the Union of the third country, territory or part thereof referred to in Boxes I.7 and I.8, or during a period where restrictive measures have been adopted by the Union against imports of these animals from this third country, territory or part thereof.
- (11) When required by the EU Member State of destination or Switzerland, in accordance with Decision 2004/558/EC and in accordance with the Agreement between the Community and the Swiss Confederation on trade in agricultural products (OJ L 114, 30.4.2002, p. 132).
- (12) Surveillance programme as laid down in Annex I to Commission Regulation (EC) No 1266/2007 (OJ L 283, 27.10.2007, p. 37).

Status: Point in time view as at 01/07/2016. **Changes to legislation:** There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model BOV-X

II.	Health information	II.a.	Certificate reference number	II.b.					
Offi	Official veterinarian								
	Name (in capital letters):		Qualification and title:						
	Date:		Signature:						
	Stamp:								

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

Model BOV-Y

COUN	TRY:					Veterinary ce	rtificate to EU	
	1.1.	Consignor	1.2.	Certificate referen	ice No	1.2.a.		
		Name Address	I.3. Central competent authority					
		Tel.	1.4.	Local competent a	authority			
ent	1.5.	Consignee	1.6.					
ignn		Name						
cons		Address						
tched		Postal code Tel.						
Part I: Details of dispatched consignment	1.7.	Country ISO of origin code origin	1.9.	Country of destination	ISO code	I.10. Region of destination	Code	
Detail	1.11.	Place of origin	1.12	<u>. </u>				
i.i.		Name Approval number Address						
a.		Address						
	I.13.	Place of loading Address Approval number	1.14	. Date of departure				
	I.15.	Means of transport	1.16	i. Entry BIP in EU				
		Aeroplane 🗆 🔁 Ship 🗖						
		Railway wagon ☐ Road vehicle ☐ Other ☐	1.17	' .				
		Identification						
	118	Documentary references Description of commodity			I 19 Commo	dity code (HS code	<u>, , , , , , , , , , , , , , , , , , , </u>	
	1.10.	Description of commounty			01.		•)	
				·		I.20. Quantity		
	1.21.					I.22. Number of	packages	
		Seal/Container No				1.24.		
	1.25.	Commodities certified for:						
		Slaughter						
1.26.				I.27. For import or	admission into	EU 🗆		
	1.28.	Identification of the commodities						
	l .	Species Breed Identi entific name)	ficatio	n system Ider	ntification numb	per Age	Sex	

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model BOV-Y

	II. I	Health information	on		II.a.	Certificate refere	nce number	II.b.			
	II.1.	Public Health	Attesta	tion							
		I, the undersign	ned offic	ial veterinarian, he	ereby c	certify, that the animals described in this certificate:					
tion		II.1.1.	last 42 6 mont	days in the case	e of br abies,	ucellosis, for the	last 30 days in the	n on health grounds, for the case of anthrax, for the last mals from holdings which did			
tifica		II.1.2.	have n	ot received:							
.ce			_	any stilbene or th	yrosta	tic substances,					
Part II: Certification			_				β - agonist substance defined in Directive 9	es for purposes other than 96/22/EC).			
		II.1.3.	with regard to bovine spongiform encephalopathy (BSE):								
		(¹) (²) either	[(a)	back to the dam	and h	erd of origin, and		n enabling them to be traced vine animals as described in) No 999/2001;			
after and g				after the date from	om whi	ch the ban on the om ruminants had	e feeding of ruminan	erned, the animals were born ts with meat-and-bone meal rced or after the date of birth ed ban.]			
		(¹) (³) or	[(a)	back to the dam	and h	erd of origin, and	,	n enabling them to be traced vine animals as described in C) No 999/2001;			
			(b)	meat-and-bone r	neal ar	nd greaves derive	d from ruminants had	he feeding of ruminants with been effectively enforced or m after the date of the feed			
		(¹) (⁴) or	[(a)	back to the dam	and h	erd of origin, and		n enabling them to be traced vine animals as described in C) No 999/2001;			
			(b)	ruminants with	meat-a ed or	ind-bone meal a after the date of b	nd greaves derived	ch the ban on the feeding of from ruminants had been indigenous case if born after			
	II.2.	Animal Health	attesta	tion:							
		I, the undersigned official veterinarian, hereby certify, that the animals described above mee requirements:						above meet the following			
		II.2.1.		ome from the territ	ory wit	h code:	(⁵) v	which, at the date of issuing			
		(1) either	[(a)	has been free for	24 m	onths from foot-an	d-mouth disease]				
		(¹) or	[(a)	without having ha	ad cas	es/outbreaks after	that date, and author	(dd/mm/yyyy), rised to export these animals (dd/mm/yyyy),]			
			(b)	has been free for 12 months from rinderpest, Rift valley fever, contagious bovine pleuropneumonia, lumpy skin disease and epizootic haemorrhagic disease, and for 6 months from vesicular stomatitis,							
			(c)	points (a) and (l) has		and imports of dom	the diseases mentioned in estic cloven-hoofed animals			
		(1) either	[(d)	has been free for	24 m	onths from blueton	gue;]				

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model BOV-Y

				_		
II.	Health informati	on		II.a.	Certificate reference number	II.b.
	(¹) or	. ,	with an inactivat against all bluet source populatio 150 km radius a	ed va ongue n as d round	24 months from bluetongue, and the accine, at least 60 days before the dayserotype/s (insert serotype/s) whemonstrated through a surveillance prothe holding(s) of origin described under the immunity period of time guarantee	ate of dispatch to the Union, ich are those present in the ogramme (9) in an area with a er box reference I.11, and the
	II.2.2.	3 month			itory described under point II.2.1 since the Union and without contact with imp	
	II.2.3.	 they have remained s under box reference I.1 			h or at least 40 days before dispatch	n in the holding(s) described
		(a)			n an area with a 150 km radius, there had a disease during the previous 60 days,	
		` ,	foot-and-mouth	diseas	n an area with a 10 km radius, there h se, rinderpest, Rift valley fever, blu py skin disease and, vesicular sto	etongue, contagious bovine
	II.2.4.				ed under a national programme for the iinst the diseases referred to in point II.	
	II.2.5.	they cor	me from herds:			
		(a)	included in an of	ficial s	ystem for the control of enzootic bovine	e leukosis, and
		(b)	that are not rest and brucellosis,		under the national legislation regardir	ng eradication of tuberculosis
		(c)	recognised as of	ficially	tuberculosis free; (6) (6a)	
	II.2.6.	they hav	ve not been vacc	inated	against brucellosis and they:	
	(1) either	[come fr	rom herds which	are red	cognised as officially brucellosis free;]	(⁶)
	(¹) or	[are cas	trated males of a	ny age	9;]	
	II.2.7.				n at least two places on their hindquariate slaughter; $(^{7})$	ters as to show that they are
	II.2.8.	they are	/were (1) dispatc	hed fro	om their holding(s) of origin, without pa	ssing through any market:
	(1) either	[directly	to the Union,]			
	(¹) or		officially authorise described under		embly centre described under box refe II.2.1]	erence I.13 situated within the
		and, unt	til dispatched to t	he Uni	on:	
					contact with other cloven-hoofed ani described in this certificate, and	mals not complying with the
		. ,			place where, or around which within e has been a case/outbreak of any o	
	II.2.9.				tainers in which they were loaded wathorised disinfectant;	vere cleaned and disinfected
	II.2.10.	they we sign of o		an offi	cial veterinarian within 24 hours of loa	ading and showed no clinical
	II.2.11.	the me	ans of transpor ted before loadin	t desc g with	atch to the Union on ribed under box reference I.15 abo an officially authorised disinfectant an flow or fall out of the vehicle or contain	ove that were cleaned and d so constructed that faeces,

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model BOV-Y

II.	Health information	II.a.	Certificate reference number	II.b.
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II.3. Animal transport attestation

I, the undersigned official veterinarian, hereby certify, that the animals described above have been treated before and at the time of loading in accordance with the relevant provisions of Regulation (EC) No 1/2005, in particular as regards watering and feeding, and they are fit for the intended transport.

Notes

This certificate is meant for live bovine animals (including Bubalus and Bison species and their cross-breeds) intended for immediate slaughter.

After importation the animals must be conveyed without delay to the slaughterhouse of destination to be slaughtered within five working days.

Part I:

— Box reference I.8:	Provide the	code	of	territory	as	appearing	in	Part	1	of	Annex I	to	Regulation	(EU)
	No 206/2010	i .												

- Box reference I.13: The assembly centre, if any, must fulfil the conditions for its approval, as laid down in Part 5 of Annex I to Regulation (EU) No 206/2010.
- Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor must inform the BIP of entry into the Union.
- Box reference I.23: For containers or boxes, the container number and the seal number (if applicable) should be included.
- Box reference I.28: Identification system: the animals must bear:

An individual number which permits tracing of their premises of origin. Specify the identification system (such as tag, tattoos, brand, chip, transponder).

An ear tag that includes the ISO code of the exporting country. The individual number must permit tracing of their premises of origin.

Species: Select amongst "Bos", "Bison" and "Bubalus" as appropriate.

Age: Date of birth (dd/mm/yyyy).

Sex (M = male, F = female, C = castrated).

Part II:

- (1) Keep as appropriate.
- (2) Only if the animals were born and continuously reared in a country or region categorised in accordance with Article 5(2) of Regulation (EC) No 999/2001 as a country or region posing a negligible BSE risk and listed as such in Decision 2007/453/EC.
- (3) Only if the country or region of origin is categorised in accordance with Article 5(2) of Regulation (EC) No 999/2001 as a country or region posing a controlled BSE risk and is listed as such in Decision 2007/453/EC.
- (4) Only if the country or region of origin has not been categorised in accordance with Article 5(2) of Regulation (EC) No 999/2001 or has been categorised as a country or region with undetermined BSE risk and is listed as such in Decision 2007/453/EC.
- (5) Code of the territory as it appears in Part 1 of Annex I to Regulation (EU) No 206/2010.
- (6) Officially tuberculosis/brucellosis free regions and herds as laid down in Annex A to Directive 64/432/EEC.
- (^{6a}) Only for a territory appearing with entry "XII" in column 6 of Part 1 of Annex I to Regulation (EU) No 206/2010 indicating that bovine herds officially declared tuberculosis-free are recognised based on equivalent conditions to those laid down in paragraphs 1 and 2 of Annex A.I to Directive 64/432/EEC for the purposes of exports to the Union of live animals certified according to the model of veterinary certificate BOV-Y.
- (7) This mark shall take the form of "L" having 13 cm in the left side and 7 cm in the bottom side with 1 cm of strength in both lines. It shall be applied using the technique known as "freeze-branding".
- (8) Date of loading. Imports of these animals shall not be allowed when the animals were loaded either prior to the date of authorisation for exportation to the Union of the third country, territory or part thereof referred to in boxes I.7 and I.8, or during a period where restrictive measures have been adopted by the Union against imports of these animals from this third country, territory or part thereof.
- (9) Surveillance programme as laid down in Annex I to Commission Regulation (EC) No 1266/2007 (OJ L 283, 27.10.2007, p. 37).

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model BOV-Y

II.	Health information	II.a.	Certificate reference number	II.b.
Official veterinarian				
	Name (in capital letters):		Qualification and title:	
	Date:		Signature:	
	Stamp:			

[F9[F10Model BOV-X-TRANSIT-RU]

Status: Point in time view as at 01/07/2016. **Changes to legislation:** There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COL	INTR				Veterinary cert	ificate to EU		
	l.1.	Consignor Name	I.2. Certificate reference	No	I.2.a.			
		Address Tel.	I.3. Central competent a	authority				
		16.	I.4. Local competent authority					
dispatched consignment	I.5.	Consignee Name Address Postal code Tel.	I.6. Person responsible f Name Address Postal code Tel.	for the load	in EU			
5	1.7.	Country of ISO code origin Russia ISO code origin Kaliningrad	I.9. Country of ISO destination Russia	O code I.	.10. Region of destination	Code		
Part I: Details	l.11.	Place of origin Name Address Postal code	1.12.					
	I.13.	Place of loading Address	I.14. Date of departure					
		Approval number						
	I.15.	Means of transport Aeroplane	I.16. Entry BIP in EU Kybartai road — Lith	huania				
			1.17.					
	I.18.	Description of commodity	I.19. Comr	modity code				
				1.20.	Quantity			
	1.21.			1.22.	Number of packages	5		
	1.23.	Seal/Container No		1.24.				
	1.25.	Commodities certified for:						
		Breeding						
	1.26.	For transit through EU to third country Third country Russian Federation ISO code RU	1.27.					
	1.28.	Identification of the commodities						
		Species Breed Identification (scientific name)	system Identific	cation numb	er Age	Sex		

Health information

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model BOV-X-TRANSIT-RU

II.a. Certificate reference No

II.b.

II.1. Animal Health attestation: I, the undersigned official veterinarian, hereby certify, that the animals described in Part I meet the following requirements: II.1.1. they come from the territory with code: RU-2 (2) which, at the date of issuing this certificate: Certification (1) either [(a) has been free for 24 months from foot-and-mouth disease;] [(a) has been considered free from foot-and-mouth disease since (1) or without having had cases/outbreaks after that date, and authorised to export these animals by Commission Implementing Regulation (EU) No, of, (dd/mm/yyyy):] Part II: (b) has been free for 12 months from rinderpest, Rift valley fever, contagious bovine pleuropneumonia, lumpy skin disease and epizootic haemorrhagic disease, and for 6 months from vesicular stomatitis; (c) where, during the last 12 months, no vaccination against the diseases referred to in points (a) and (b) has been carried out and imports of domestic cloven-hoofed animals vaccinated against these diseases are not permitted; (1) either [(d) has been free for 24 months from bluetongue;] [(d) has not been free for 24 months from bluetongue, and the animals have been vaccinated with an inactivated vaccine, at least 60 days before the date of the movement, against all bluetongue serotype/s (insert serotype/s) which are those present in the source population as demonstrated through a surveillance programme (4) in an area with a 150 km radius around the holding(s) of origin described under box reference I.11., and the animals are still within the immunity period of time guaranteed in the specifications of the vaccine;] (1) either origin are kept:1 (1) or [II.1.2. they have remained in the territory with code RU-2 since birth, or for at least the last six months before the date of dispatch via the European Union and without contact with imported cloven-hoofed animals for the last 30 days;] II.1.3. they have remained [since birth or at least 40 days before the date of dispatch (5) in the holding(s) of origin described under box reference I.11.: (a) in and around which, in an area with a 150 km radius, there has been no case/outbreak of epizootic haemorrhagic disease during the previous 60 days: (b) in and around which, in an area with a 10 km radius, there has been no case/outbreak of foot-and-mouth disease, rinderpest, Rift valley fever, bluetongue, contagious bovine pleuropneumonia, lumpy skin disease and vesicular stomatitis during the previous 40 days; II.1.4. they are not animals to be killed under a national programme for the eradication of diseases, nor have they been vaccinated against the diseases referred to under point II.1.1., (a) and (b), and: (a) they did not come in contact with other cloven-hoofed animals not complying with the health requirements as described in this certificate: (b) they were not at any place where, or around which, within a 10 km radius, during the previous 30 days there has been a case/outbreak of any of the diseases referred to in point II.1.1.; II.1.5. any transport vehicles or containers in which they were loaded were cleaned and disinfected before loading with an officially authorised disinfectant: II.1.6. they were examined by an official veterinarian within 24 hours of loading and showed no clinical sign of disease; II.1.7. they have been loaded for dispatch to Russia via the European Union on .. (dd/mm/yyyy) (3) in the means of transport described under box reference I.15. above that were cleaned and disinfected before loading with an officially authorised disinfectant and so constructed that faeces, urine, litter or fodder could not flow or fall out of the vehicle or container during transportation; II.1.8. the consignment is intended to leave the European Union at the designated Border Inspection Post Medininkai, Lithuania

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY		Model BOV-X-TRANSIT-R		
II. Health information	II.a. Certificate reference No	II.b.		
II.2. Animal transport attestation				
I, the undersigned official veterinarian, hereby certify, that th loading in accordance with the relevant provisions of Council and they are fit for the intended transport.				
Notes:				
This certificate is meant for transit through the European Union of domes breeds) intended for breeding and/or production coming from the region				
Part I:				
- Box reference I.8.: Provide the code of territory as appearing in Part	t 1 of Annex I to Commission Regulation	on (EU) No 206/2010.		
 Box reference I.13.: The assembly centre, if any, must fulfil the cond Regulation (EU) No 206/2010. 	ditions for its approval, as laid down in	Part 5 of Annex I to Commission		
 Box reference I.15.: Registration number of road vehicle is to be pro Border Inspection Post of entry into the Union. 	vided. In case an emergency, the con-	signor must immediately inform the		
— Box reference I.23.: For containers or boxes, the container number a	and the seal number (if applicable) mu	st be included.		
Box reference I.28.: Identification system: the animals must bear:				
 An individual number which permits tracing of their premises of c transponder). 	origin. Specify the identification system	(such as tag, tattoos, brand, chip		
- An ear tag that includes the ISO code of the exporting country.	. The individual number must permit	tracing of their premises of origin		
 Box reference I.28.: Species: select amongst "Bos", "Bison" and "Bul 	balus" as appropriate.			
 Box reference I.28.: Age: date of birth (dd/mm/yy). 				
— Box reference I.28.: Sex (M = male, F = female, C = castrated).				
— Box reference I.28.: Breed: select purebred, cross-breed.				
Part II:				
(¹) Keep as appropriate.				
$(^2)$ Code of the territory as it appears in Part 1 of Annex I to Commission	ion Regulation (EU) No 206/2010.			
3) Date of loading. Transit of these animals shall not be allowed when the animals were loaded either prior to the date of authorisation for transit to Russia via the European Union from this third country, territory or part thereof referred to in Boxes I.7., or during a period where restrictive measures have been adopted by the European Union against transit of these animals from this third country, territory or part thereof via the European Union.				
(4) Surveillance programme as laid down in Annex I to Commission Re	gulation (EC) No 1266/2007.			
$(^5)$ Delete the text in square brackets if the second option for point II.1.	.2. is deleted.			
Official veterinarian/Official inspector				
Name (in capital letters):	Qualification	tion and title:		
Date:	Signature	e:		

[F11Model OVI-X]]

Stamp:

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

cou	INTR	1		Veterinary certificate to EU
	l.1.	Consignor Name	I.2. Certificate reference No	1.2.a.
		Address Tel.	I.3. Central competent authorit	ty
ent		Tel.	I.4. Local competent authority	
signm	1.5.	Consignee Name	1.6.	
dispatched consignment		Address Postal code Tel.		
Part I: Details of disp	1.7.	Country of origin ISO code I.8. Region of origin Code	I.9. Country of ISO cod destination	e I.10. Region of Code destination
Deta	l.11.	Place of origin	I.12.	
Part I:		Name Approval number Address		
	I.13.	Place of loading	I.14. Date of departure	
		Address Approval number		
	I.15.	Means of transport	I.16. Entry BIP in EU	
		Aeroplane Ship Railway wagon Road vehicle Other O		
		Identification Documentary references	1.17.	
	I.18.	Description of commodity	I.19. Commodity	code (HS code)
				I.20. Quantity
	I.21.			I.22. Number of packages
	1.23.	Seal/Container No		1.24.
	1.25.	Commodities certified for:		
		Breeding	Fattening	
	1.26.		I.27. For import or admission in	nto EU 🔲
	1.28.	Identification of the commodities	•	
		Species Breed Identification (scientific name) system	Identification number	Age Sex

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model OVI-X II. Health information II.a. Certificate reference number II.b. II.1. **Public Health Attestation** I, the undersigned official veterinarian, hereby certify, that the animals described in this certificate: II.1.1. come from holdings which have been free from any official prohibition on health grounds, for the last 42 days in the case of brucellosis, for the last 30 days in the case of anthrax, for the last six months in the case of rabies, and, have not been in contact with animals from holdings which did not comply with these conditions; Certification II.1.2. have not received any stilbene or thyrostatic substances, oestrogenic, androgenic, gestagenic or β- agonist substances for purposes other than therapeutic or zootechnic treatment (as defined in Directive 96/22/EC). 11.2. Animal Health attestation Part I, the undersigned official veterinarian, hereby certify, that the animals described above meet the following requirements: II.2.1. they come from the territory with code: (1), which, at the date of issuing this certificate: (2) either [(a) has been free for 24 months from foot-and-mouth disease,] (2) or [(a) has been considered free from foot-and-mouth disease since (dd/mm/yyyy), (dd/mm/yyyy),] (b) has been free for 12 months from rinderpest, Rift valley fever, peste des petits ruminants, sheep pox and goat pox, contagious caprine pleuropneumonia, and epizootic haemorrhagic disease, and for 6 months from vesicular stomatitis, (c) where during the last 12 months, no vaccination against the diseases mentioned in points (a) and (b) has been carried out and imports of domestic cloven-hoofed animals vaccinated against these diseases are not permitted;] (2) either [(d) has been free for 24 months from bluetongue;] [(d) has been free for 24 months from bluetongue, and the animals have reacted negatively to a serological test for the detection of antibody for bluetongue and epizootic haemorrhagic disease, carried out on two occasions on samples of blood taken at the beginning of the isolation/quarantine period and at least 28 days later, (2)(7) or (dd/mm/yyyy) and on (dd/mm/yyyy), the second of which must have been taken within 10 days before export;] (d) has not been free for 24 months from bluetonque, and the animals have been vaccinated with an inactivated vaccine, at (2) or has not been free for 24 months from bluetongue, and the animals have been vaccinated with an inactivated vaccine, at least 60 days before the date of dispatch to the Union, against all bluetongue serotype/s. .. (insert serotype/s) which are those present in the source population as demonstrated through a surveillance programme (9) in an area with a 150 km radius around the holding(s) of origin described under box reference I.11., and the animals are still within the immunity period of time guaranteed in the specifications of the vaccine;] II.2.2. they have remained in the territory described under point II.2.1. since birth, or for at least the last six months before dispatch to the Union and without contact with imported cloven-hoofed animals for the last 30 days; II.2.3. they have remained since birth or at least 40 days in the holding(s) described under box reference I.11. before dispatch (a) in and around which, in an area with a 150 km radius, there has been no case/outbreak of epizootic haemorrhagic disease during the previous 60 days, and (b) in and around which, in an area with a 10 km radius, there has been no case/outbreak of foot-and-mouth disease, rinderpest, Rift valley fever, bluetongue, peste des petits ruminants, sheep pox and goat pox, contagious caprine pleurop-neumonia and vesicular stomatitis during the previous 40 days; II.2.4. according to my knowledge and to the written declaration made by the owner, the animals: (a) do not come from holdings, and have not been in contact with animals of a holding, in which the following diseases have (i) contagious agalactia of sheep or goats (Mycoplasma agalactiae, Mycoplasma capricolum, Mycoplasma mycoides var. mycoides large colony), within the last six months, (ii) paratuberculosis and caseous lymphadenitis, within the last 12 months, (iii) pulmonary adenomatosis, within the last three years, and (iv) Maedi/Visna or caprine viral arthritis/encephalitis: (2) either [within the last three years.] [within the last 12 months, and all the infected animals were slaughtered and the remaining animals subsequently reacted negatively to two tests carried out at least six months apart.]

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model OVI-X Health information II.a. Certificate reference number II.b. (b) are included in an official system for notification of these diseases, and (c) have been free from clinical or other evidence of tuberculosis and brucellosis during the three years prior to export; they are not animals to be killed under a national programme for the eradication of diseases, nor have they been vaccinated against the diseases referred to in point II.2.1.(a) and (b); 11.2.5. II.2.6. they originate: (2)(3) either [from the territory described under box reference I.8., which has been recognised as officially brucellosis-free;] (2) or [from the holding(s) described under box reference I.11., where, in respect of brucellosis (Brucella melitensis): (a) all susceptible animals have been free from clinical or any signs of this disease for the last 12 months, (b) a representative number of the domestic ovine and caprine animals over an age of six months are submitted each year to a serological test, (4)] (²)(⁵) either [(c) all domestic ovine or caprine animals have not been vaccinated against this disease, save those vaccinated with Rev. 1 vaccine more than two years ago; (d) the last two tests (6), separated by an interval of at least six months, carried out on (dd/mm/yyyy) on all domestic ovine and caprine animals over six months of age gave negative results, and] [(c) domestic ovine or caprine animals under the age of 7 months are vaccinated against this disease with Rev. 1 (2) or (d) the last two tests (6), separated by an interval of at least six months, carried out: on ... (dd/mm/yyyy) on all non-vaccinated domestic ovine and caprine animals over six months of (dd/mm/yyyy) and on (dd/mm/yyyy) on all vaccinated domestic and on ovine and caprine animals over 18 months of age gave negative results, and] (e) there are only domestic ovine and caprine animals that comply with the above conditions and requirements;] the uncastrated rams have been kept continuously during the previous 60 days in a holding where no case of contagious epididymitis (*Brucella ovis*) has been diagnosed in the last 12 months and, these rams have undergone during the previous 30 days a complement fixation test to detect contagious epididymitis with a result of less than 50 IU/ml;] (2) [II.2.7. 11.2.8 they have been kept continuously since birth in a country where the following conditions are fulfilled: (a) classical scrapie is compulsorily notifiable; (b) an awareness, surveillance and monitoring system for classical scrapie is in place; (c) ovine and caprine animals affected with classical scrapie are killed and completely destroyed; (d) 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, and [II.2.8.1 they are animals intended for production and they are destined for a Member State other than those with a negligible risk status for classical scrapie approved in accordance with point 2.2 of Section A of Chapter A of Annex VIII to Regulation (EC) No 999/2001, or other than those which are listed in point 3.2 of Section A of Chapter A of Annex VIII to Regulation (EC) (2) either No 999/2001 as having an approved national scrapie control programme;] [II.2.8.1 they are animals intended for breeding and they are destined for a Member State other than those with a negligible risk status for classical scrapie approved in accordance with point 2.2 of section A of chapter A of Annex VIII to Regulation (EC) No 999/2001, or other than those which are listed in point 3.2 of Section A of Chapter A of Annex VIII to Regulation (EC) No 999/2001 as having an approved national scrapie control programme and: (2) or [they come from a holding or holdings that have complied with the requirements laid down in point 1.3 of Section A of Chapter A of Annex VIII to Regulation (EC) No 999/2001;]] (2) either [they are ovine animals of the ARR/ARR prion protein genotype and they come from a holding where no official movement restriction has been imposed due to BSE or classical scrapie during the last two years;]] (2) or

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model OVI-X Health information II.a. Certificate reference number II.b. (2) or they are destined for a Member State with a negligible risk status for classical scrapie approved in accordance with point 2.2 of Section A of Chapter A of Annex VIII to Regulation (EC) No 999/2001, or for a Member State listed in point 3.2 of Section A of Chapter A of Annex VIII to Regulation (EC) No 999/2001 as having an approved national scrapie control programme, III.2.8.1 [they come from a holding or holdings that have complied with the requirements laid down in point 1.2 of Section A of Chapter A of Annex VIII to Regulation (EC) No 999/2001;]] (2) either [they are ovine animals of the ARR/ARR prion protein genotype and they come from a holding where no official movement restriction has been imposed due to BSE or classical scrapie during the last two years;]] (2) or II.2.9. they are/were (2) dispatched from their holding(s) of origin, without passing through any market, (2) either [directly to the Union,] [to the officially authorised assembly centre described under box reference I.13. situated within the territory described under point II.2.1.,] (2) or and, until dispatched to the Union: (a) they did not come in contact with other cloven-hoofed animals not complying with the health requirements as described in this certificate, and (b) they were not at any place where, or around which within a 10 km radius, during the previous 30 days there has been a case/outbreak of any of the diseases referred to in point II.2.1.; II.2.10. any transport vehicles or containers in which they were loaded were cleaned and disinfected before loading with an officially authorised disinfectant; II.2.11. they were examined by an official veterinarian within 24 hours of loading and showed no clinical sign of disease; so constructed that faeces, urine, litter or fodder could not flow or fall out of the vehicle or container during transportation Animal transport attestation I, the undersigned official veterinarian, hereby certify, that the animals described above have been treated before and at the time of loading in accordance with the relevant provisions of Regulation (EC) No 1/2005, in particular as regards watering and feeding, and they are fit for the intended transport. This certificate is meant for live domestic ovine animals (Ovis aries) and domestic caprine animals (Capra hircus) intended for breeding or After importation the animals must be conveyed without delay to the holding of destination where they shall remain for a minimum period of 30 days before further movement outside the holding, except in the case of a dispatch to a slaughterhouse Box reference I.8.: Provide the code of territory as appearing in Part 1 of Annex I to Regulation (EU) No 206/2010. - Box reference I.13.: The assembly centre, if any, must comply with the conditions for its approval, as laid down in Part 5 of Annex I to Regulation (EU) No 206/2010. Box reference I.15.: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In
case of unloading and reloading, the consignor must inform the BIP of entry into the Union. Box reference I.19.: Use the appropriate HS code: 01.04.10 or 01.04.20. - Box reference I.23.: For containers or boxes, the container number and the seal number (if applicable) should be included.

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

cou	NTRY			Model OVI-)	
П.	Health infor	mation	II.a. Certificate reference number	II.b.	
— E	Box reference I.28.:	Identification system: The animals must bear:			
		An individual number which permits tracing of tattoos, brand, chip, transponder) and the anato		identification system (such as tag,	
		An ear tag that includes the ISO code of the export origin.	orting country. The individual number n	nust permit tracing of their premises	
		Species: Select amongst "Ovis aries" and "Capi	ra hircus" as appropriate.		
		Age: (months).			
		Sex (M = male, F = female, C = castrated).			
Part	t II:				
(1) (Code of the territory	as it appears in Part 1 of Annex I to Regulation	n (EU) No 206/2010.		
(²) ł	Keep as appropriate).			
(3) (Only for a territory a	appearing with the entry "V" in column 6 of Part	1 of Annex I to Regulation (EU) No 20	06/2010.	
) a	The representative number of animals to be tested for brucellosis must, for each holding, consist of: all non-castrated male animals, which have not been vaccinated against brucellosis, over six months old, all non-castrated male animals, which have been vaccinated against brucellosis, over 18 months old, all animals brought onto the holding since the previous tests, and 25% of females which are sexually mature, within a minimum of 50 females.				
(⁵) 1	This must be comple	eted when the destination is a Member State or pa	art of a Member State listed in one of the	he Annexes of Decision 93/52/EEC.	
		Part 6 of Annex I to Regulation (EU) No 206/201 ne holding of origin is involved the date of the m		be clearly indicated.	
		antees to be provided when required in column 5 ongue and for Epizootic-haemorrhagic-disease in			
)´ e	exportation to the U	ports of these animals shall not be allowed whi Inion of the third country, territory or part thereon adopted by the Union against imports of these	of referred to in boxes I.7. and I.8., o	or during a period where restrictive	
(⁹) 8	Surveillance progran	nme as laid down in Annex I to Commission Reg	gulation (EC) No 1266/2007 (OJ L 28	3, 27.10.2007, p. 37).	
Offic	cial veterinarian				
	Name (in capital le	etters):	Qualification a	and title:	
	Date:		Signature:		
	Stamp:				

Status: Point in time view as at 01/07/2016. **Changes to legislation:** There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

Model OVI-Y

COL	JNTR	Υ	Veterinary certificate to EU
	1.1.	Consignor Name	I.2. Certificate reference No I.2.a.
		Address	I.3. Central competent authority
		Tel.	
nent			I.4. Local competent authority
signr	1.5.	Consignee	1.6.
cons		Name Address	
ped		Postal code	
patc		Tel.	
Part I: Details of dispatched consignment	1.7.	Country of ISO code I.8. Region of Code origin Code	I.9. Country of ISO code I.10. Region of destination Code
etail		Discount white	
<u>:</u>	1.11.	Place of origin	1.12.
Part		Name Approval number Address	
		Addiese	
	I.13.	Place of loading	I.14. Date of departure
		Address Approval number	
	I.15.	Means of transport	I.16. Entry BIP in EU
		Aeroplane	
		Road vehicle Other	I.17.
		Identification	
	110	Documentary references	110. Commodity and (HS ands)
	1.18.	Description of commodity	I.19. Commodity code (HS code)
			I.20. Quantity
	1.21.		I.22. Number of packages
	122	Seal/Container No	1.24.
			1.24.
	1.25.	Commodities certified for:	
		Slaughter	
	1.26.		I.27. For import or admission into EU
	1.28.	Identification of the commodities	
		Species Breed Identification (scientific name) system	Identification number Age Sex

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model OVI-Y II. Health information II.a. Certificate reference number II.b. II.1. **Public Health Attestation** I, the undersigned official veterinarian, hereby certify, that the animals described in this certificate: II.1.1. come from holdings which have been free from any official prohibition on health grounds, for the last 42 days in the case of brucellosis, for the last 30 days in the case of anthrax, for the last six months in the case of rabies, and, have not been in contact with animals from holdings which did not satisfy these conditions; II: Certification II.1.2. have not received: Part - any stilbene or thyrostatic substances, — oestrogenic, androgenic, gestagenic or β- agonist substances for purposes other than therapeutic or zootechnic treatment (as defined in Directive 96/22/EC). 11.2. Animal Health attestation I, the undersigned official veterinarian, hereby certify, that the animals described above meet the following requirements: II.2.1. they come from the territory with code:(1) which, at the date of issuing this certificate: (2) either [(a) has been free for 24 months from foot-and-mouth disease] (2) or (b) has been free for 12 months from rinderpest, Rift valley fever, peste des petits ruminants, sheep pox and goat pox, contagious caprine pleuropneumonia, and epizootic haemorrhagic disease, and for 6 months from vesicular stomatitis (c) where during the last 12 months, no vaccination against the diseases mentioned in points (a) and (b) has been carried out and imports of domestic cloven-hoofed animals vaccinated against these diseases are not permitted; (2) either [(d) has been free for 24 months from bluetongue;] (2) or (d) has not been free for 24 months from bluetongue, and the animals have been vaccinated with an inactivated has not been tree for 24 months from billetongue, and the animals have been vaccinated with an inactivated vaccine, at least 60 days before the date of dispatch to the Union, against all bluetongue serotype/s. which are those present in the source population as demonstrated through a surveillance programme (5) in an area with a 150 km radius around the holding(s) of origin described under box reference I.11., and the animals are still within the immunity period of time guaranteed in the specifications of the vaccine;] II.2.2. they have remained in the territory described under point II.2.1. since birth, or for at least the last three months before dispatch to the Union and without contact with imported cloven-hoofed animals for the last 30 days; II.2.3. they have remained since birth or at least 40 days before dispatch in the holding(s) described under box reference I.11: (a) in and around which in an area with a 150 km radius there has been no case/outbreak of epizootic haemorrhagic disease during the previous 60 days, and (b) in and around which, in an area with a 10 km radius, there has been no case/outbreak of foot-and-mouth disease, rinderpest, Rift valley fever, bluetongue, peste des petits ruminants, sheep pox and goat pox; contagious caprine pleuropneumonia and vesicular stomatitis during the previous 40 days; II.2.4. they are not animals to be killed under a national programme for the eradication of diseases, nor have they been vaccinated against the diseases referred to in point II.2.1(a) and (b); II.2.5. they are/were (2) dispatched from their holding(s) of origin, without passing through any market, (2) either [directly to the Union]

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model OVI-Y

Health information II.a. Certificate reference number II.b. [to the officially authorised assembly centre described under box reference I.13 situated within the territory described under point II.2.1,] and, until dispatched to the Union: (a) they did not come in contact with other cloven-hoofed animals not complying with the health requirements as described in this certificate, and (b) they were not at any place where, or around which within a 10 km radius, during the previous 30 days there has been a case/outbreak of any of the diseases referred to in point II.2.1; II.2.6. in respect of scrapie: (2) (3) [II.2.6.1. if they are destined for a Member State which benefits, for all or part of its territory, from the provisions laid down in points (b) or (c) of Chapter A(I) of Annex VIII to Regulation (EC) No 999/2001, the animals comply with the guarantees provided for in the programmes referred to in those points, as laid down in Article 2 of Regulation (EC) 546/2006, and] (2) either [II.2.6.2. were born in and continuously reared on holdings in which a case of scrapie has never been diagnosed;] [II.2.6.2. are domestic ovine animals of the ARR/ARR prion protein genotype as defined in Annex I to Decision 2002/1003/EC, coming from a holding where no case of scrapie has been reported in the last six months;] (2) or 11.2.7. any transport vehicles or containers in which they were loaded were cleaned and disinfected before loading with an officially authorised disinfectant; they were examined by an official veterinarian within 24 hours of loading and showed no clinical sign of disease; they have been loaded for dispatch to the Union on ... (dd/mm/yyyy) (4) in the means of transport described under box reference I.15 above that were cleaned and disinfected before loading with an officially authorised disinfectant and so constructed that faeces, urine, litter or fodder could not flow or fall out of the vehicle or container during transportation. Animal welfare attestation II.3. I, the undersigned official veterinarian, hereby certify, that the animals described above have been treated before and at the time of loading in accordance with the relevant provisions of Regulation (EC) No 1/2005, in particular as regards watering and feeding, and they are fit for the intended transport. Notes This certificate is meant for live domestic ovine animals (Ovis aries) and domestic caprine animals (Capra hircus) intended for immediate slaughter After importation the animals must be conveyed without delay to the slaughterhouse of destination to be slaughtered within five working days. Part I: Box reference I.8: Provide the code of territory as appearing in Part 1 of Annex I to Regulation (EU) No 206/2010. Box reference I.13: The assembly centre, if any, must fulfil the conditions for its approval, as laid down in Part 5 of Annex I to Regulation (EU) No 206/2010. - Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor must inform the BIP of entry into the Union. Box reference I.19: Use the appropriate HS code: 01.04.10 or 01.04.20. - Box reference I.23: For containers or boxes, the container number and the seal number (if applicable) should be included.

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

co	UNTRY		Model OVI-Y			
II.	Health information	II.a. Certificate reference number	II.b.			
F	Box reference I.28: Identification system: The animals must bear:					
	 An individual number which permits tracing of their premises of origin. Specify the identification system (such as tag, tattoos, brand, chip, transponder) and the anatomic place used in the animal. 					
	- An ear tag that includes the ISO code of the exporting country. The individual number must permit tracing of their premises of origin.					
	Species: Select amongst "Ovis aries" and "Capra hircus" as appropri	iate.				
	Age: months.					
	Sex (M = male, F = female, C = castrated).					
Pa	rt II:					
(1)) Code of the territory as it appears in Part 1 of Annex I to Regulation (EU) No 206/2010.					
(2)	Keep as appropriate.					
(³)	Guarantees in relation to a programme of control of scrapie, as requ and Chapter E of Annex IX to Regulation (EC) No 999/2001.	ested by the EU Member State of des	tination, in application of Article 15			
(⁴)	Date of loading. Imports of these animals shall not be allowed whe exportation to the Union of the third country, territory or part there measures have been adopted by the Union against imports of these	of referred to in boxes I.7 and I.8, or	r during a period where restrictive			
(⁵)	Surveillance programme as laid down in Annex I to Commission Reg	gulation (EC) No 1266/2007 (OJ L 283	3, 27.10.2007, p. 37.).			
Of	ficial veterinarian					
	Name (in capital letters):	Qualification and title:				
	Date:	Signature:				
	Stamp:					

[F8Model POR-X]

Status: Point in time view as at 01/07/2016. **Changes to legislation:** There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Veterinary certific							
	l.1.	Consignor Name Address	I.2. Certificate reference No I.2.a.				
		Tel.	I.3. Central competent authority				
ment			I.4. Local competent authority				
consign	1.5.	Consignee Name	1.6.				
Part I: Details of dispatched consignment		Address Postal code Tel.					
of di							
: Details	1.7.	Country ISO I.8. Region Code of origin	I.9. Country of destination code I.10. Region of destination	Code			
art	l.11.	Place of origin	1.12.				
_		Name Approval number Address					
	I.13.	Place of loading	I.14. Date of departure				
		Address Approval number	·				
	I.15.	Means of transport	I.16. Entry BIP in EU				
		Aeroplane Ship Railway wagon CRoad vehicle Other Uldentification	l.17.				
		Documentary references					
	I.18.	Description of commodity	I.19. Commodity code (HS code) 01.03				
			I.20. Quantity				
	I.21.		I.22. Number of packag	es			
	1.23.	Identification of container/seal number	1.24.				
	I.25. Commodities certified for:						
		Breeding					
	1.26.		I.27. For import or admission into EU				
	I.28. Identification of the commodities						
		Species Identification system Identification number Age (scientific name)					

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model POR-X

II. Health information II.a. Certificate reference number II.b. II.1. **Public Health Attestation** I, the undersigned official veterinarian, hereby certify, that the animals described in this certificate: II.1.1. come from holdings which have been free from any official prohibition on health grounds, for the last 42 days in the case of brucellosis, for the last 30 days in the case of anthrax and for the past six months in the case of rabies and, the animals have not been in contact with animals from holdings which did not satisfy these conditions; Part II: Certification II.1.2. have not received: - any stilbene or thyrostatic substances — oestrogenic, androgenic, gestagenic or β-agonist substances for purposes other than therapeutic or zootechnic treatment (as defined in Directive 96/22/EC). • (1) (2) (10) [II.1.3. are domestic porcine animals either coming from a holding officially recognised as applying controlled housing conditions in accordance with Article 8 of Regulation (EC) No 2075/2005 or are not weaned and less than 5 weeks of age.] 11.2 Animal Health attestation I, the undersigned official veterinarian, hereby certify, that the animals described above meet the following requirements: II.2.1. they come from the territory with code: (1) which, at the date of issuing this certificate (²) either [(a) has been free for 24 months from foot-and-mouth disease, for 12 months from rinderpest, African swine fever classical swine fever, swine vesicular disease and vesicular exanthema, and] [(a) (i) has been free [for 24 months from foot-and-mouth disease] (²), for 12 months from rinderpest, African swine fever, vesicular exanthema, [classical swine fever] (²) and [swine vesicular disease] (²), and (2) or (2) either (b) for 6 months from vesicular stomatitis, and [(b) the animals have been kept for the 21 days, or since birth if younger than 21 days of age, prior to entering the pre-export quarantine in a holding in which no case of vesicular stomatitis was officially reported during that period and (2) (9) or during the pre-export quarantine of not less than 30 days prior to shipment in a quarantine station protected from vector insects where they were subjected with negative results at a serum dilution of 1 in 32 to a virus neutralisation test for vesicular stomatitis carried out as referred to in Part 6 of Annex I to Regulation (EU) No 206/2010 on samples taken at least 21 days after commencement of the quarantine; and] (c) where during the last 12 months, no vaccination against these diseases has been carried out and imports of domestic cloven-hoofed animals vaccinated against these diseases are not permitted; II.2.2. they have remained in the territory described under point II.2.1 since birth, or for at least the last six months before dispatch to the Union and without contact with imported cloven-hoofed animals for the last 30 days; II.2.3. they have remained in the holding(s) described under box reference I.11 since birth, or for at least 40 days prior to dispatch and, during this period, in the holding(s) and in an area with a 10 km radius around the holding(s) of origin, there has been no case/outbreak of the diseases referred to in point II.2.1; II.2.4. A they are not animals to be killed under a national programme for the eradication of diseases, nor have they been vaccinated (2) (3) [II.2.4. B they have been subjected within the past 30 days to a test for swine vesicular disease antibodies and a test for classical swine fever antibodies with negative results in both cases;] (2) (4) [II.2.4. C they have been subjected within the past 30 days to a buffered Brucella antigen test for porcine brucellosis with negative II.2.5 they come from herds which are not restricted under the national brucellosis eradication programme; II.2.6 they are/were (2) dispatched from their holding(s) of origin, without passing through any market, (2) either [directly to the Union,]

[to the officially authorised assembly centre described under box reference I.13 situated within the territory described under

point II.2.1,]

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY				Model POR-X
II.	Health information	II.a. Certificate reference number	II.b.	

II.a. Certificate reference number and, until dispatched to the Union: (a) they did not come in contact with other cloven-hoofed animals not complying with the health requirements as described in (b) they were not at any place where, or around which within a 10 km radius, during the previous 40 days there has been a case/outbreak of any of the diseases referred to in point II.2.1, and (c) in the case the country has not been free for 6 months of vesicular stomatitis, they were transported to the place of loading protected from vector insects; II.2.7. any transport vehicles or containers in which they were loaded were cleaned and disinfected before loading with an officially authorised disinfectant; II.2.8. they were examined by an official veterinarian within 24 hours of loading and showed no clinical sign of disease; II.3. Animal transport attestation I, the undersigned official veterinarian, hereby certify, that the animals described above have been treated before and at the time of loading in accordance with the relevant provisions of Regulation (EC) No 1/2005, in particular as regards watering and feeding, and they are fit for the intended transport. (2) (6) [II.4. Specific requirements II.4.1. Aujeszky's disease is notifiable in the country referred to in box reference I.7; II.4.2. according to official information, no clinical, pathological or serological evidence of Aujeszky's disease has been recorded for the last 12 months in the holding(s) of origin referred to in box reference I.11., and in those holdings situated in its vicinity within 5 km: II.4.3. the animals referred to in box reference I.28: (a) prior to dispatch for exportation, have remained since birth in the holding(s) of origin referred to in box reference I.11. or they have remained in this(ese) holdings(s) for the last 3 months and in others of equivalent status since birth, (b) have been isolated in accommodation approved by the competent authority for the last 30 days immediately prior to dispatch for export, without direct or indirect contact with other Suidae animals, (c) have been subjected to an ELISA test for the presence of Ig (7) on sera taken at least 21 days after entry into isolation, with negative results; and, all animals in isolation have also given negative results to this test, and (d) have not been vaccinated against Aujeszky's disease and have not been in contact with vaccinated animals and the herd of origin has not been vaccinated during the previous 12 months.] (2) (8) [II.4.4. (further requirements and/or tests) This certificate is meant for live domestic porcine animals (Sus scrofa) intended for breeding or production. After importation the animals must be conveyed without delay to the holding of destination where they shall remain for a minimum period of 30 days before further movement outside the holding, except in the case of animals dispatched directly to a slaughterhouse or of animals transiting the Union from one third country to another third country.

Part I

- Box reference I.8: Provide the code of territory as appearing in Part 1 of Annex I to Regulation (EU) No 206/2010.
- Box reference I.13: The assembly centre, if any, must fulfil the conditions for its approval, as laid down in Part 5 of Annex I to Regulation (EU)
 No. 206/2010

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

II.	Health information	II.a. Certificate reference number	II.b.				
	Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. case of unloading and reloading, the consignor must inform the BIP of entry into the Union.						
	— Box reference I.23: For containers or boxes, the container number and the seal number (if applicable) should be included.						
	Box reference I.28.: Identification system: the animals must bear:						
	 An individual number which permits tracing of their premises of transponder). 	origin. Specify the identification system	(such as tag, tattoos, brand, chip,				
	- An ear tag that includes the ISO code of the exporting country. The individual number must permit tracing of their premises of						
— Box reference I.28: Age: months.							
	Box reference I.28.: Sex (M = male, F = female, C = castrated).						
-	Part II:						
	(1) Code of the territory as it appears in Part 1 of Annex I to Regulation (EU) No 206/2010.						
	(²) Keep as appropriate.						
•	3) Supplementary guarantees to be provided when required in column entry 'B'.	nn 5 'SG' of Part 1 of Annex I to Regu	ulation (EU) No 206/2010, with the				
•	4) Supplementary guarantees to be provided when required in columentry 'C'.	nn 5 'SG' of Part 1 of Annex I to Regu	ulation (EU) No 206/2010, with the				
•	5) Date of loading. Imports of these animals shall not be allowed w exportation to the Union of the third country, territory or part there measures have been adopted by the Union against imports of thes	eof referred to in boxes I.7. and I.8., o	r during a period where restrictive				
•	⁶) When required by the EU Member State of destination or Switzerlar the Community and the Swiss Confederation on trade in agricultural in column 6 'Specific conditions' of Part 1 of Annex I to Regulation	products (OJ L 114, 30.4.2002, p. 132)					
•	7) To be carried out according to the standards laid down in Annex III used shall be the whole virus ELISA.	to Decision 2008/185/EC. In the case of	f pigs aged over 4 months, the test				
	8) Further requirements requested by Finland in respect of transmissil	ble gastro-enteritis.					
	9) Supplementary guarantees to be provided when required in columentry 'D'.	n 5 'SG' of Part 1 of Annex I to Regu	ulation (EU) No 206/2010, with the				
· ⁽¹⁾ (Only for third countries with the entry 'XI' in column 6 'Specific co	onditions' in Part 1 of Annex I to Regul	ation (EU) No 206/2010. ◀				
-	Official veterinarian						
	Name (in capital letters):	Qualifica	tion and title:				
	Date:	Signature	e:				
	Stamp:						

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

Model POR-Y

	со	UNTRY						Veterinary ce	rtificate to EU	
	1.1.	I.1. Consignor			I.2. Certific	ate referenc	e numbe	r I.2.a.		
		Name			I.3. Central Competent Authority					
	Address Tel. No									
				I.4. Local Competent Authority						
ıt	I.5. Consignee		I.6.							
umei		Name								
ısig	Address Postal code									
Part I: Details of dispatched consignment										
		Tel. No								
	1.7.	Country ISO code	I.8. Region of origin	Code	I.9. Countr destina		ISO code	I.10. Region of destination	Code	
	1.11.	. Place of origin			l.12.					
Deta		Name	Approval number							
1:1		Address								
Pai		Name Address	Approval number				/			
		Name Address	Approval number							
	I.13	. Place of loading Address	Approval number		I.14. Date of	f departure		time of departure		
	I.15. Means of transport			I.16. Entry BIP in EU						
	Aeroplane Ship Railway wagon									
	Road vehicle Other			147						
	Identification:			1.17.						
		Documentary references:								
	I.18. Description of commodity					I.19. Com	modity c	ode (HS code)	01.03	
						I.20. Quantity				
	1.21.					I.22. Number of packages				
	I.23. Identification of container/seal number					1.24.				
	1.25	. Commodities certified fo	r:							
		Slaughter								
	1.26.			I.27. For import or admission into EU						
	1.28	I.28. Identification of the commodities								
		Species Identification (Scientific name) system			Identification number		А	ge	Sex	

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model POR-Y

Health information II.a. Certificate reference number II.b. II.1. **Public Health Attestation** I, the undersigned official veterinarian, hereby certify, that the animals described in this certificate: come from holdings which have been free from any official prohibition on health grounds, for the last 42 days in the case of brucellosis, for the last 30 days in the case of anthrax and for the past six months in the case of rabies and, the animals have not been in contact with animals from holdings which did not satisfy these conditions; Part II: Certification II.1.2 have not received: - any stilbene or thyrostatic substances, oestrogenic, androgenic, gestagenic or β- agonist substances for purposes other than therapeutic or zootechnic treatment (as defined in Directive 96/22/EC). ▶⁽¹⁾ (²)(⁵)[II.1.3 are domestic porcine animals either coming from a holding officially recognised as applying controlled housing conditions in accordance with Article 8 of Regulation (EC) No 2075/2005 or are not weaned and less than 5 weeks of 11.2. **Animal Health attestation** I, the undersigned official veterinarian, hereby certify, that the animals described above meet the following requirements: II.2.1 they come from the territory with code:(1) which, at the date of issuing this certificate: [(a) has been free for 24 months from foot-and-mouth disease, for 12 months from rinderpest, African (2) either swine fever, classical swine fever, swine vesicular disease and vesicular exanthema, and for 6 months from vesicular stomatitis, and] [(a) (i) has been free [for 24 months from foot-and-mouth disease] (2), for 12 months from rinderpest, (2) or African swine fever, vesicular exanthema, [classical swine fever] (2) and [swine vesicular disease] (2), and for 6 months from vesicular stomatitis, and (ii) has been considered free from [foot-and-mouth disease] (2), [classical swine fever] (2) and [swine vesicular disease] (2), since (dd/mm/yyyy), without having had cases/outbreaks from that date, and authorised to export these animals by Commission Regulation (EU) No, of (dd/mm/yyyy), and] (b) where during the last 12 months, no vaccination against these diseases has been carried out and imports of domestic cloven-hoofed animals vaccinated against these diseases are not they have remained in the territory described under point II.2.1 since birth, or for at least the last three months before 11.2.2 dispatch to the Union and without contact with imported cloven-hoofed animals for the last 30 days; 11.2.3 they have remained in the holding(s) described under box reference I.11 since birth, or for at least 40 days prior to dispatch, and, during this period, in the holding(s) and in an area with a 10 km radius around the holding(s) of origin, there has been no case/outbreak of the diseases referred to in point II.2.1; they are not animals to be killed under a national programme for the eradication of diseases, nor have they been vaccinated against the diseases referred to in point II.2.1; II.2.5 they are/were (2) dispatched from their holding(s) of origin, without passing through any market, (2) either [directly to the Union,] [to the officially authorised assembly centre described under box reference I.13 situated within the (2) or territory described under point II.2.1,] and, until dispatched to the Union: (a) they did not come in contact with other cloven-hoofed animals not complying with the health requirements as (b) they were not at any place where, or around which within a 10 km radius, during the previous 40 days there has been a case/outbreak of any of the diseases referred to in point II.2.1;

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model POR-Y

II. Health information II.a. Certificate reference number II.b.

- II.2.6 any transport vehicles or containers in which they were loaded were cleaned and disinfected before loading with an officially authorised disinfectant;
- II.2.7 they were examined by an official veterinarian within 24 hours of loading and showed no clinical sign of disease;

II.3. Animal transport attestation

I, the undersigned official veterinarian, hereby certify, that the animals described above have been treated before and at the time of loading in accordance with the relevant provisions of Regulation (EC) No 1/2005, in particular as regards watering and feeding, and they are fit for the intended transport.

(2) (4) [II.4. Specific requirements

- II.4.1 Aujeszky's disease is notifiable in the country referred to in box reference I.7;
- II.4.2 according to official information, no clinical, pathological or serological evidence of Aujeszky's disease has been recorded in the holding(s) of origin referred to in box reference I.11, for the last 3 months;
- II.4.3 the animals referred to in box reference I.28:
 - (a) have remained in the holding(s) of origin referred to in box reference I.11 since birth or for the last 60 days prior to dispatch for exportation, and
 - (b) have not been vaccinated against Aujeszky's disease.]

Notes

This certificate is meant for live domestic porcine animals (Sus scrofa) intended for immediate slaughter after importation.

After importation the animals must be conveyed without delay to the slaughterhouse of destination to be slaughtered within five working days.

Part I:

- Box reference I.8: Provide the code of territory as appearing in Part 1 of Annex I to Regulation (EU) No 206/2010.
- Box reference I.13: The assembly centre, if any, must fulfil the conditions for its approval, as laid down in Part 5 of Annex I to Regulation (EU) No 206/2010.
- Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor must inform the BIP of entry into the Union.
- Box reference I.23: For containers or boxes, the container number and the seal number (if applicable) should be included.
- Box reference I.28: Identification system: The animals must bear:
 - An individual number which permits tracing of their premises of origin. Specify the identification system (such as tag, tattoos, brand, chip, transponder) and the anatomic place used in the animal.
 - An ear tag that includes the ISO code of the exporting country. The individual number must permit tracing of their premises of origin.
- Box reference I.28: Age: months.
- Box reference I.28: Sex (M = male, F = female, C = castrated).

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model POR-Y

II.	Health information	II.a. Certificate reference number	II.b.					
Pa	rt II:							
(¹)	Code of the territory as it appears in Par	rt 1 of Annex I to Regulation (EU) No 206/2010	ı.					
(2)	Keep as appropriate.							
(3)	(3) Date of loading. Imports of these animals shall not be allowed when the animals were loaded either prior to the date of authorisation for exportation to the Union of the third country, territory or part thereof referred to in boxes I.7 and I.8, or during a period where restrictive measures have been adopted by the Union against imports of these animals from this third country, territory or part thereof.							
(4)	When required by the EU Member State	e of destination, in accordance with Decision 2	008/185/EC.					
▶ (1)(5)	Only for third countries with the entry 'X	I' in column 6 'Specific conditions' in Part 1 of	Annex I to Regulation (EU) No 206/2010. ◀					
Off	icial veterinarian							
	Name (in capital letters):	Qualification	and title:					
	Date:	Signature:						
		<u> </u>						
	Stamp:							

Status: Point in time view as at 01/07/2016. **Changes to legislation:** There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

Model RUM

COL	COUNTRY Veterinary certificate to EU									
	l.1.	Consignor Name	I.2. Certificate reference No I.2.a.							
		Address Tel.	I.3. Central competent authority							
뒽			I.4. Local competent authority							
nsignme	1.5.	Consignee Name Address	1.6.							
Part I: Details of dispatched consignment		Postal code Tel.								
ls of disp	1.7.	Country of origin ISO code I.8. Region of origin Code	I.9. Country of destination ISO code l.10. Region of destination Code							
Detai	l.11.	Place of origin	1.12.							
Part I:		Name Approval number Address								
	I.13.	Place of loading	I.14. Date of departure							
		Address Approval number								
	l.15.	Means of transport	I.16. Entry BIP in EU							
		Aeroplane Ship Railway wagon Road vehicle Other Ship								
		Identification Documentary references	I.17. No(s) of CITES							
	I.18.	Description of commodity	I.19. Commodity code (HS code)							
			I.20. Quantity							
	I.21.		I.22. Number of packages							
	1.23.	Seal/Container No	1.24.							
	1.25.	Commodities certified for:								
		Breeding	Slaughter							
	1.26.		I.27. For import or admission into EU							
	1.28.	Identification of the commodities								
		Species Identification system Identific (scientific name)	eation number Age Sex							

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model RUM II. Health information II.a. Certificate reference number II.b. II.1. **Public Health Attestation** I, the undersigned official veterinarian, hereby certify, that the animals described in this certificate: II.1.1. come from a holding which has been free from any official prohibition on health grounds, for the last 42 days in the case of brucellosis and tuberculosis, for the last 30 days in the case of anthrax, for the last six months in the case of rabies, and, have not been in contact with animals from holdings which did not satisfy these conditions; Part II: Certification - any stilbene or thyrostatic substances, oestrogenic, androgenic, gestagenic or β - agonist substances for purposes other than therapeutic or zootechnic treatment (as defined in Directive 96/22/EC). 11.2. Animal Health Attestation I, the undersigned official veterinarian, hereby certify, that the animals described above meet the following requirements: II.2.1. they come from the territory with code: (1) which, at the date of issuing this certificate: (a) has been free for 24 months from foot-and-mouth disease and bluetongue, for 12 months from rinderpest, Rift valley fever, contagious bovine pleuropneumonia, lumpy skin disease, peste des petits ruminants, sheep pox and goat pox, contagious caprine pleuropneumonia and epizootic haemorrhagic disease and for six months from vesicular stomatitis, and (b) where during the last 12 months, no vaccination against foot-and-mouth disease, rinderpest, Rift valley fever, contagious bovine pleuropneumonia, lumpy skin disease, peste des petits ruminants, sheep pox and goat pox, contagious caprine pleuropneumonia and epizootic haemorrhagic disease and during the last 24 months no vaccination against bluetongue has been carried out and imports of cloven-hoofed animals vaccinated against these diseases are not permitted; II.2.2. they have remained [in the territory described under point II.2.1. since birth, or for at least the last six months before dispatch to the (2) either Union and without contact with cloven-hoofed animals imported into this territory less than six months ago;] [in the country of dispatch for at least 60 days since entry, if they are animals of the relevant species listed in Part 7 of Annex I to Regulation (EU) No 206/2010 and they were imported directly under the conditions specified for each species in Part 7 of Annex I to Regulation (EU) No 206/2010 from a third country during a period of less than six months prior to embarkation to the Union and in any case they have been separated from other animals (2) or not of the same health status after being released in the exporting country and before exportation to the II.2.3. they have remained since birth or at least 40 days before dispatch in the holding/establishment (2) described under boxes reference I.11 and I.13: (a) in and around which in an area of radius of 150 km, there has been no case/outbreak of bluetongue and epizootic haemorrhagic disease during the previous 60 days, and (b) in and around which in an area of 10 km radius, there has been no case/outbreak of the other diseases referred to in point II.2.1 during the previous 40 days; II.2.4. they are not animals to be killed under a national programme for the eradication of diseases, nor have they been vaccinated against any of the diseases referred to in point II.2.1, and they: (2) (4) either [come from a herd which is recognised as officially tuberculosis free, and] (2) (5) or [have been subjected to an intradermal tuberculin test within the past 30 days with negative results, and] they have not been vaccinated against brucellosis and they: (2) (4) either [come from a herd which is recognised as officially brucellosis free;] (2) (5) or [have been subjected to a serum agglutination test which showed a brucella count of less than 30 IU of agglutination per ml, within the past 30 days;] (2) or [are castrated males of any age:]

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

П.	Health	information II.a. Certificate reference number II.b.
	7100.0	
	II.2.5.	according to my knowledge and to the written declaration made by the owner, the animals:
		 (a) do not come from holdings/establishments (2), and have not been in contact with animals of a holding/establishment, in which the following diseases have been clinically detected:
		 (i) contagious agalactia of sheep or goats (Mycoplasma agalactiae, Mycoplasma capricolum, Mycoplasma mycoides va mycoides 'large colony'), within the last six months,
		(ii) paratuberculosis and caseous lymphadenitis, within the last 12 months,
		(iii) pulmonary adenomatosis, within the last three years, and
		(iv) Maedi/Visna or caprine viral arthritis/encephalitis,
		(2) either [within the last three years,]
		(2) or [within the last 12 months, and all the infected animals were slaughtered and the remaining animals subsequently reacted negatively to two tests carried out at least six months apart,]
		(b) are included in an official system for notification of these diseases, and
		(c) have been free from clinical or other evidence of tuberculosis and brucellosis during the three years prior to export;
(²) (⁶) [II.2.6.	the animals have reacted negatively to a serological test for the detection of antibody for bluetongue and epizootic-haemo rhagic-disease, carried out on two occasions on samples of blood taken at the beginning of the isolation/quarantine period an at least 28 days later on
	II.2.7.	they are dispatched from the holding/establishment described under boxes reference I.11 and I.13 directly to the Union and, undispatched to the Union:
		(a) they did not come in contact with other cloven-hoofed animals not complying with the health requirements as described this certificate, and
		(b) they were not at any place where, or around which within a 10 km radius, during the previous 30 days there has been case/outbreak of any of the diseases referred to in point II.2.1;
	II.2.8.	any transport vehicles or containers in which they were loaded were cleaned and disinfected before loading with an official authorised disinfectant;
	II.2.9.	they were examined by an official veterinarian within 24 hours of loading and showed no clinical sign of disease;
	II.2.10	they have been loaded for dispatch to the Union on
II.3.	Anima	I transport attestation
	loading	undersigned official veterinarian, hereby certify, that the animals described above have been treated before and at the time of a lacordance with the relevant provisions of Regulation (EC) No 1/2005, in particular as regards watering and feeding, and the for the intended transport.
(²) (⁸)	[II.4. Specif	ic requirements
	II.4.1.	According to official information, no clinical or pathological evidence of infectious bovine rhinotracheitis (IBR) has been recorde in the holding/establishment (²) of origin referred to in boxes reference I.11 and I.13, for the last 12 months;
	II.4.2.	the animals referred to in box reference I.28.:
		(a) have been isolated in accommodation approved by the competent authority for the last 30 days immediately prior to dispatch for export, and

(b) have been subjected to a serological test for IBR on sera taken at least 21 days after entry into isolation, with negative results, and all animals in isolation have also given negative results to this test, and

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model RUM II.a. Certificate reference number II.b. Health information (c) have not been vaccinated against IBR.; (²) [II.4.3. (further requirements and/or tests) Notes This certificate is meant for live animals of the order Artiodactyla (excluding bovine animals (including *Bubalus* and *Bison* species and their cross-breeds), Ovis aries, Capra hircus, Suidae and Tayassuidae), and of the families Rhinocerotidae and Elephantidae. Use one certificate per species. After importation the animals must be conveyed without delay to the holding of destination where they shall remain for a minimum period of 30 days before further movement outside the holding, except in the case of a dispatch to a slaughterhouse Part I: Box reference I.8.: Provide the code of territory as appearing in Part 1 of Annex I to Regulation (EU) No 206/2010. Box reference I.13.: The assembly centre, if any, must fulfil the conditions for its approval, as laid down in Part 5 of Annex I to Regulation (EU) No 206/2010. Box reference I.15.: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In
case of unloading and reloading, the consignor must inform the BIP of entry into the Union. Box reference I.19.: Use the appropriate HS code: 01.02, 01.04.10, 01.04.20 or 01.06.19. - Box reference I.23.: For containers or boxes, the container number and the seal number (if applicable) should be included. Box reference I.28.: Identification system: Specify the identification system (tag, tattoos, brand, chip, transponder). The ear tag includes the ISO code of the exporting country. The individual number must permit tracing of their premises of origin. Sex (M = male, F = female, C = castrated). Species: Select the species amongst those listed for the following families: Antilocapridae: Antilocapra spp.: Addax spp., Aepyceros spp., Alcelaphus spp., Ammodoroas spp., Ammotragus spp., Antidoroas spp., Antilope spp., Bose-laphus spp., Budoroas spp., Capra spp. (excluding Capra hircus), Cephalophus spp., Connochaetes spp., Damaliscus spp. (including Beatragus), Doroatragus spp., Gazella spp., Hemitragus spp., Hippotragus spp., Kobus spp., Litocranius spp., Madoqua spp., Naemorhedus spp. (including Nemorhaedus and Capricomis), Botragus spp., Oreamnos spp., Oreotragus spp., Oryx spp., Ourebia spp., Ovibos spp., Ovis spp. (excluding Ovis aries), Pantholops spp., Pelea spp., Procapra spp., Pseudois spp., Pseudoryx spp., Raphicerus spp., Redunca spp., Rupicapra spp., Salga spp., Signoceros-Alecelaphus spp., Subvicars spp., Spp., Salga spp., Signoceros-Alecelaphus spp., Tragelaphus spp., Signoceros-Alecelaphus spp., Tragelaphus spp., Signoceros-Alecelaphus spp., Tragelaphus spp., Signoceros-Alecelaphus spp., Tragelaphus spp., Signoceros-Alecelaphus spp., Sp Bovidae: Sylvicapra spp., Syncerus spp., Taurotragus spp., Tetracerus spp., Tragelaphus spp. (including Boocerus). Camelidae: Camelus spp., Lama spp., Vicugna spp. Alces spp., Axis-Hyelaphus spp., Blastocerus spp., Capreolus spp., Cervus-Rucervus spp., Dama spp., Elaphurus spp., Hippocamelus spp., Hydropotes spp., Mazama spp., Megamuntiacus spp., Muntiacus spp., Odocoileus spp., Ozotoceros Cervidae: spp., Pudu spp., Rangifer spp. Giraffidae: Giraffa spp., Okapia spp. Hippopotamidae: Hexaprotodon-Choeropsis spp., Hippopotamus spp., Moschidae: Moschus spp. Tragulidae: Hvemoschus spp., Tragulus-Moschiola spp., Rhinocerotidae: Ceratotherium spp., Dicerorhinus spp., Diceros spp., Rhinoceros spp. Elephantidae: Elephas spp., Loxodonta spp., as appropriate.

COUNTRY

Model RUM

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

П.	Health information	II.a. Certificate reference number	11.6.						
Pa	Part II:								
(¹)	(1) Code of the territory as it appears in Part 1 of Annex I to Regulation (EU) No 206/2010.								
(²)	Keep as appropriate.								
(3)	3) In this case the health certificate has to be accompanied by the official document on quarantine and test conditions laid down in Part 2 of Annex I to Regulation (EU) No 206/2010 (model "CAM").								
(4)	b) Officially tuberculosis/brucellosis free regions or herds recognised as equivalent to the requirements laid down in Annex A to Directive 64/432/EEC and which appear in column 6 of Part 1 of Annex I to Regulation (EU) No 206/2010, with the entry "VII" as regards tuberculosis, "VIII", as regards brucellosis.								
(⁵)	5) Tests carried out in accordance with the protocols that, for the disease concerned, are described in Part 6 of Annex I to Regulation (EU) No 206/2010. However for the tuberculin test a result of an increase in skin fold thickness of 2mm or more, or clinical signs of such as oedema, exudation, necrosis, pain and/or inflammation shall be deemed to be positive.								
(⁶)	Supplementary guarantees to be provided when required in column 5 "A". Tests for Bluetongue and for Epizootic-haemorrhagic-disease i								
(7)	7) Date of loading. Imports of these animals shall not be allowed when the animals were loaded either prior to the date of authorisation for exportation to the Union of the third country, territory or part thereof referred to in boxes I.7. and I.8., or during a period where restrictive measures have been adopted by the Union against imports of these animals from this third country, territory or part thereof.								
(8)	When required by the EU Member State of destination.								
Off	icial veterinarian								
	Name (in capital letters):	Qualification and t	itle:						
	Date:	Signature:							
	Stamp:								

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

Model SUI

	co	UNTRY					Veterinary ce	rtificate to EU
	l.1.	Consignor		I.2. Certification	ate reference	number	I.2.a.	
		Name		I.3. Central Competent Authority				
		Address		-				
		Tel. No		I.4. Local C	ompetent Aut	thority		
nt	I.5.	Consignee	I.6.					
nme		Name						
nsig		Address						
Ор		Postal code						
che		Tel. No						
Part I: Details of dispatched consignment	1.7.	Country ISO I.8. Region Coforigin code of origin	Code	I.9. Country destina		SO ode	I.10. Region of destination	Code
ils o	1.11.	. Place of origin		I.12.				
Deta		Name Approval number						
1::		Address						
Pa		Name Approval number Address						
		Name Approval number Address						
	1.13	. Place of loading Address Approval number		I.14. Date of	departure	tir	me of departure	
	1.15	. Means of transport		I.16. Entry B	IP in EU			
		Aeroplane Ship Railway wagon						
		Road vehicle Other		147 N. () . (CITEC			
		Identification: Documentary references:		I.17. No(s) of CITES				
	I.18	. Description of commodity		I.19. Commodity code (HS code)				
						I.20. Q	uantity	
	1.21					I.22. N	umber of packag	jes
	1 23	s. Identification of container/seal number				1.24.		
	1.20	. Northingalion of container south tamber				1.24.		
	1.25	. Commodities certified for:						
		Breeding Fat	tening			Slauç	ghter	
	1.26			I.27. For imp	ort or admiss	ion into E	U	
	1.28	. Identification of the commodities						
		Species Identification (Scientific name) system		Identification number	1	Age	Э	Sex

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model SUI

	II.	Health	information	II.a. Certificate reference number	II.b.	
tion	II.1.	Public	Health Attestation			
		I, the u	indersigned official veterina	arian, hereby certify, that the animals describe	d in this certificate:	
		II.1.1	case of brucellosis, for th	ch has been free from any official prohibition on the last 30 days in the case of anthrax and for the thin in contact with animals from holdings which	e past six months in th	e case of rabies and,
rtifica		II.1.2	have not received:			
Part II: Certification			 any stilbene or thyros 	static substances,		
Part				enic, gestagenic or β - agonist substances for particle of the Directive 96/22/EC).	urposes other than ther	apeutic or zootechnic
	II.2.	Anima	l Health attestation			
		I, the u	indersigned official veterina	arian, hereby certify, that the animals describe	d above meet the follow	ving requirements:
		II.2.1	they come from the territor	ory with code: (1) which	n, at the date of issuing	this certificate:
				months from foot-and-mouth disease, for 12 r r, swine vesicular disease and vesicular exa		
				t 12 months, no vaccination against these dis lls vaccinated against these diseases are not p		ed out and imports of
		II.2.2		e territory described under point II.2.1 since bi I without contact with cloven-hoofed animals in		
		II.2.3	dispatch, and, during this	e holding described under boxes reference I.1 e period, in the holding(s) and in an area with a outbreak of the diseases referred to in point II.2	10 km radius around th	
		II.2.4 A	vaccinated against the di	be killed under a national programme for the esseases referred to in point II.2.1 and they have test for porcine brucellosis with negative resu	been subjected within	
	(²) (³) [II.2.4 B		ed within the past 30 days to a test for swine bodies with negative results in both cases]	vesicular disease anti	bodies and a test for
	(2) (4) [II.2.4 C	they have been subjecte negative results]	d within the past 30 days to a buffered Bruce	ella antigen test for po	rcine brucellosis with
		11.2.5	they come from holdings	which:		
				nder a national control and eradication progeschen disease), and	ramme for brucellosis	s, porcine enteroviral
			(b) are included in an off	ficial system for notification of these diseases;		
		II.2.6	they are dispatched from dispatched to the Union:	the holding described under boxes reference	I.11 and I.13 directly to	o the Union and, until
			(a) they did not come in described in this cert	contact with other cloven-hoofed animals not ificate, and	complying with the he	ealth requirements as
				place where, or around which within a 10 km rake of any of the diseases referred to in point II.2		ous 40 days there has

Health information

II.

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model SUI

II.b.

II.a. Certificate reference number

	II.2.7 any transport vehicles or containers in which they were loaded were cleaned and disinfected before loading with officially authorised disinfectant;							
	II.2.8	they were examined by an official veterinarian within 24 hours of loading and showed no clinical sign of disease;						
	II.2.9	transport described under	or dispatch to the Union on box reference I.15 above that were cleaned that faeces, urine, litransportation.	ed and disinfected before loading with an				
II.3.	Animal	transport attestation						
	I, the undersigned official veterinarian, hereby certify, that the animals described above have been treated before and at the time of loading in accordance with the relevant provisions of Regulation (EC) No 1/2005, in particular as regards watering and feeding, and they are fit for the intended transport.							
(²) (6) [II.4.	. Specifi	c requirements						
	II.4.1	Aujeszky's disease is notifia	able in the country referred to in box reference	ee I.7;				
	II.4.2		nation, no clinical, pathological or serologica nths in the holding(s) of origin referred to in b the holding(s);					
	II.4.3	the animals referred to in be	ox reference I.28:					
			exportation, have remained since birth in or they have remained in this holding for the					
			accommodation approved by the competen port, without direct or indirect contact with other.					
			o an ELISA test for the presence of gl antib n negative results; and, all animals in isolation					
			ed against Aujeszky's disease and have not l oot been vaccinated during the previous 12 m					
(2) (8)	[11.4.4]]	(further requirements and/or tests)				

Notes

This certificate is meant for live non-domestic Suidae (*Babyrousa* spp., *Hylochoerus* spp., *Phacochoerus* spp., *Potamochoerus* spp., and *Sus* spp.), Tayassuidae (*Catagonus* spp., *Pecari* spp., *Tayassu* spp.) and Tapiridae (*Tapirus* spp.).

After importation the animals must be conveyed without delay to the holding of destination where they shall remain for a minimum period of 30 days before further movement outside the holding, except in the case of a dispatch to a slaughterhouse.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

Model SUI

CC	UNINI		wodel Sc					
II.	Health information	II.a. Certificate reference number	II.b.					
Pa	rt I:							
_ _	 Box reference I.8: Provide the code of territory as appearing in Part 1 of Annex I to Regulation (EU) No 206/2010. Box reference I.13: The assembly centre, if any, must fulfil the conditions for its approval, as laid down in Part 5 of Annex I to Regulation (EU) No 206/2010. 							
_		r (railway wagons or container and lorries), fli ading, the consignor must inform the BIP of er						
—	Box reference I.19: Use the appropriate	HS code: 01.03 or 01.06.19.						
—	Box reference I.23: For containers or bo	xes, the container number and the seal numb	er (if applicable) should be included.					
—	Box reference I.28: Identification system	r: The animals must bear:						
	 An individual number which permits brand, chip, transponder) and the ar 	s tracing of their premises of origin. Specify th natomic place used in the animal.	e identification system (such as tag, tattoos,					
	 An ear tag that includes the ISO coo origin. 	de of the exporting country. The individual nur	mber must permit tracing of their premises of					
—	Box reference I.28: Age: months.							
—	Box reference I.28: Sex (M = male, F = f	emale, C = castrated).						
—	Box reference I.28: Species.							
Pa	rt II:							
(¹)	Code of the territory as it appears in Par	rt 1 of Annex I to Regulation (EU) No 206/2010	0.					
(²)	Keep as appropriate.							
(3)	Supplementary guarantees to be provide with the entry 'B'.	ded when required in column 5 'SG' of Part 1	of Annex I to Regulation (EU) No 206/2010,					
(4)	Supplementary guarantees to be provide with the entry 'C'.	ded when required in column 5 'SG' of Part 1	of Annex I to Regulation (EU) No 206/2010,					
(5)	for exportation to the Union of the third	s shall not be allowed when the animals were I country, territory or part thereof referred to ir d by the Union against imports of Suidae an	n boxes I.7 and I.8, or during a period where					
(⁶)	When required by the EU Member State	e of destination, in accordance with Decision 2	2008/185/EC.					
(7)	To be carried out according to the stan 4 months, the test used shall be the who	dards laid down in Annex III to Decision 200 ole virus ELISA.	8/185/EC. In the case of animals aged over					
(8)	Further requirements requested by Finla	and in respect of transmissible gastro-enteritis	S.					
Off	icial veterinarian							
	Name (in capital letters):	Qualification	n and title:					
	Date:	Signature:						
	Stamp:							

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

Model CAM Specific animal health attestation for animals quarantined in St. Pierre and Miquelon prior to introduction into the Union

	CO	UNTRY	•				Veterinary cert	ificate to EU
	1.1.	Consignor		I.2. Certific	ate reference i	number	I.2.a.	
		Name		I.3. Central Competent Authority				
		Address						
		Tel. No		I.4. Local C	ompetent Auti	nority		
ŧ	1.5.	Consignee		I.6.				
nme		Name						
nsig		Address				/		
o p		Postal code						
che		Tel. No						
Part I: Details of dispatched consignment	1.7.	Country ISO of origin code	I.8. Region Code of origin	I.9. Country destina		SO I.	.10. Region of destination	Code
ils o	1.11.	. Place of origin		I.12.				
Deta		Name	Approval number					
± ::		Address						
Ра		Name Address	Approval number					
		Name	Approval number					
		Address	Approvarnumber					
	1.13.	. Place of loading		I.14. Date of departure time of departure				
		Address	Approval number					
	I.15.	. Means of transport Aeroplane Shi	p	I.16. Entry BIP in EU				
		Road vehicle Othe	er 🗌	I.17. No(s) of CITES				
		Identification:		1.17. NO(5) OF CITES				
		Documentary references:			I			
	I.18	. Description of commodity			I.19. Commo	odity cod	e (HS code)	01.06.19
						1.20. Qu	uantity	
	1.21					1.22. Nu	ımber of packages	5
	1.23	3. Identification of container/se	eal number			1.24.		
	1.25	i. Commodities certified for:						
		Breeding			Slaug	hter		
	1.26	5.	I.27. For imp	ort or admissi	on into El	U		
	1.28	B. Identification of the commod	dities					
		Species (Scientific name)	Identification system	Identification number	n	Age	•	Sex

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model CAM

	000					model oran			
	II.	Health	information		II.a. Certificate reference number	II.b.			
ication	II.1.	Quarantine conditions attestation							
		(date (date)	dd/mm/yyyy) of of Annex I to Reg	released entry (²)) julation (E period the	arian, hereby certify, that the animals describe on	been resident fromquelon under the conditions provided for in before being released for exportation to the			
Part II: Certification		II.1.1.	Brucellosis:						
Part			(a) B. abortus: least 42 day		gglutination Test (SAT) and Rose Bengal Test (F	RBT) within two days after arrival and after at			
			(b) B. ovis: Cor	nplement	Fixation Test (CFT) within two days after arriva	al and after at least 42 days			
			(c) B. melitensi	is: SAT an	d RBT within two days after arrival and after at	least 42 days			
		II.1.2.	Bluetongue and	d Epizooti	c haemorrhagic disease				
			(5) either	[two test 21 days]	s using Bluetongue competitive Elisa test wit	hin two days after arrival and after at least			
			(⁵) or		ve been quarantined for more than 60 days and free of Bluetongue vectors (<i>Culicoides</i>), and J.				
		II.1.3.	Tuberculosis						
					lin test according to annex B to Directive 64, s after arrival and after at least 42 days from the				
		II.1.4.	Foot-and-mouth after arrival and		: ELISA test for the detection of antibodies areast 42 days	d a virus neutralizaton test within two days			
		II.1.5.	Rinderpest: cor	mpetitive E	ELISA test within two days after arrival and after	er at least 42 days			
		II.1.6.	Vesicular stoma	atitis: ELIS	SA or virus- neutralisation test within two days a	after arrival and after at least 42 days			
		II.1.7.	Rift valley fever	: an ELIS	A test or a virus neutralisation test within two da	ays after arrival and after at least 42 days			
		II.1.8.	Lumpy skin dise	ease: ELIS	SA or virus neutralisation test within two days a	fter arrival and after at least 42 days			
		II.1.9.	Crimean Congo haemorrhagic fever: ELISA or virus neutralisation test within two days after arrival and after at leas 42 days						
		II.1.10.	Surra: blood mi	croscopy	within two days after arrival and after at least 4	2 days			
		II.1.11.	Malignant catar	rhal fever	: immunofluorescence test within two days afte	er arrival and after at least 42 days			
	II.2.	Supple	ementary guara	ntees					
		II.2.1	Bovine leukosis Member State o		st or ELISA within two days after arrival and afte tion) $(^5)$	er at least 42 days (When required by the EU			

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model CAM

II.	Health information			II.a. Certificate reference number		II.b.			
II.3.	Treatm	reatments							
	They h	ney have been subjected to:							
	II.3.1.	3.1. an internal and external antiparasitic treatment during the quarantine period							
	II.3.2.								
		(5) either	[a treatm	ent with streptomycin 25mg/kg]					
		(5) or	-	iotic treatment effective against Le	eptospira sp	pp. (specify			
	(⁵) [II.3.3.	3. a vaccination against rabies (if requested) on							

Notes

This certificate is meant for live animals of the family Camelidae.

Part I:

- Box reference I.8: Provide the code of territory as appearing in Part 1 of Annex I to Regulation (EU) No 206/2010.
- Box reference I.13: The assembly centre, if any, must fulfil the conditions for its approval, as laid down in Part 5 of Annex I to Regulation (EU) No 206/2010.
- Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor must inform the BIP of entry into the Union.
- Box reference I.23: For containers or boxes, the container number and the seal number (if applicable) should be included.
- Box reference I.28: Identification system: The animals must bear:
 - An individual number which permits tracing of their premises of origin. Specify the identification system (such as tag, tattoos, brand, chip, transponder) and the anatomic place used in the animal.
 - An ear tag that includes the ISO code of the exporting country. The individual number must permit tracing of their premises of origin.
- Box reference I.28: Age: months.
- Box reference I.28: Sex (M = male, F = female, C = castrated).
- Box reference I.28: Species: Select amongst 'Camelus spp.', 'Lama spp.', 'Vicugna spp.' as appropriate.

Part II:

- (¹) Animal health certificate for non domestic animals other than Suidae, consigned to the Union (model 'RUM') as laid down in Part 2 of Annex I to Regulation (EU) No 206/2010.
- (2) Date in which the last animal in a group entered the quarantine facility.
- (3) Tests performed in accordance with the methods described in Chapter 2 of Part 7 of Annex I to Regulation (EU) No 206/2010.
- (4) Results of the tests performed must be attached in original to this health attestation.
- (5) Keep as appropriate.
- NB: Sampling and testing procedures must be grouped as much as possible while respecting the minimum time intervals to avoid excessive handling and manipulation of the animals.

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNT	HY		Model CAN
II.	Health information	II.a. Certificate reference number	II.b.
Official	veterinarian		
	Name (in capital letters):	Qualification	and title:
	Date:	Signature:	
	Stamp		

PART 3

Addendum for transport of animals by sea

(To be completed and attached to the veterinary certificate when transport to the Union frontier includes, even for part of the journey, transportation by ship.)

Declaration by the master of the ship

I, the undersigned, master of ship (name ...), declare that the animals referred to in the attached veterinary certificate No ... have remained on board the ship during the voyage from ... in ... (exporting country) to ... in the Union and that the ship did not call at any place outside ... (exporting country) en route to the Union other than: ... (Ports of call en route). Moreover, during the journey, these animals have not been in contact with other animals on board of a lower health status.

Done at ... on ...

(Port of arrival)	(Date of arrival)		
(stamp)	(signature of master)		
	(name in capital letters and title)		

PART 4

Addendum for transport of animals by air

(To be completed and attached to the veterinary certificate when transport to the Union frontier includes, even for part of the journey, transportation by air.)

Declaration by the captain of the aircraft

I, the undersigned, captain of the aircraft (name ...), declare that the crate or container and the area around the crate or container containing the animals referred to in the attached veterinary certificate No ... has been sprayed with insecticide before departure. Done at ... on ...

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

(Airport of departure)	(Date of departure)			
(stamp)	(signature of captain)			
	(name in capital letters and title)			

PART 5

Conditions for the approval of assembly centres (referred to in Article 4)

In order to be approved, assembly centres must meet the following requirements:

- I. They must be supervised by an official veterinarian.
- II. They must each be situated at the centre of an area of at least 20 km in diameter in which, according to official findings, there has been no case of foot-and-mouth disease for at least a period of 30 days prior to their use as approved assembly centres.
- III. They must, before each use as approved assembly centres, be cleansed and disinfected with a disinfectant officially authorised in the exporting country as effective for the control of foot-and-mouth disease.
- IV. They must have, taking into account their animal capacity:
 - (a) a facility dedicated exclusively for use as an assembly centre;
 - (b) appropriate facilities, that are easy to clean and disinfect, for loading, unloading and adequate housing of a suitable standard for the animals, for watering and feeding them, and for giving them any necessary treatment;
 - (c) appropriate facilities for inspection and isolation;
 - (d) appropriate equipment for cleaning and disinfecting rooms and trucks;
 - (e) an appropriate storage area for fodder, litter and manure;
 - (f) an appropriate system for collecting and disposal of waste water;
 - (g) an office for the official veterinarian.
- V. When operating, they must have sufficient veterinarians to carry out all duties set out in Part 5;
- VI. They must only admit animals that are individually identified so as to guarantee traceability. To this end, when animals are admitted the owner or the person in charge of the centre must ensure that the animals are properly identified and accompanied by health documents or certificates for the species and categories involved.

In addition, the owner or the person in charge of the assembly centre must record on a register or in a data base, and retain for at least three years the name of the owner, the origin of the animals, the dates of entry and exit, the identification number of the animals or registration number of the herd of origin and the holding of destination, and, the registration number of the carrier and the registration number of the lorry delivering or collecting animals from that assembly centre.

VII. All animals passing through the assembly centre must fulfil the health conditions established for the introduction of the relevant category of animal into the Union.

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the
Commission Regulation (EU) No 206/2010. (See end of Document for details)

- VIII. Animals to be introduced into the Union which pass through an assembly centre must, within six days of arrival at the assembly centre, be loaded and dispatched directly to the border of the exporting country:
 - (a) without coming into contact with cloven-hoofed animals other than animals which fulfil the health conditions established for the introduction of the relevant category of animal into the Union;
 - (b) segregated into consignments so that no consignment contains both animals for breeding or production and animals for immediate slaughter;
 - (c) in transport vehicles or containers which have first been cleansed and disinfected with a disinfectant officially authorised in the exporting country as effective for the control of foot-and-mouth disease and which are so constructed that faeces, urine, litter or fodder cannot flow or fall out during transportation.
- IX. Where the conditions for the export of animals to the Union require that a test is carried out within a specified period before loading, that period must include any period of assembly, up to six days, from the date of arrival of the animals at the approved assembly centre.
- X. The exporting third country must designate the centres which are approved for animals for breeding and production and those centres which are approved for animals for slaughter and must notify the Commission and the competent central authorities of the Member States of the names and addresses of such premises. That information must be updated regularly.
- XI. The exporting third country shall determine the procedure for official supervision of approved assembly centres and shall ensure that such supervision is carried out.
- XII. The approved assembly centres must be regularly inspected by the competent authority of the third country in order to check that the requirements for approval set out in points I to XI continue to be fulfilled.

If those inspections show that those conditions are no longer complied with, the approval of the centre must be suspended. The approval may be restored only when the competent authority of the third country is satisfied that the centre fully complies with the conditions set out in points I to XI.

PART 6

Protocols for the standardisation of materials and testing procedures

(referred to in Article 5)

Tuberculosis (TBL)

The single intradermal tuberculin test using bovine tuberculin shall be carried out according to Annex B to Directive 64/432/EEC. In the case of Suidae animals, the single intradermal tuberculin test using avian tuberculin shall be carried out according to Annex B to 64/432/EEC, except that the site of injection shall be the loose skin at the base of the ear.

I^{F12}Brucellosis (Brucella abortus) (BRL)

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

The serum agglutination test, complement fixation test, buffered brucella antigen test, enzymelinked immunosorbent assays (ELISA) and fluorescence polarisation assay (FPA) shall be carried out according to Annex C to Directive 64/432/EEC.] Brucellosis (Brucella melitensis) (BRL)

Tests shall be carried out according to Annex C to Directive 91/68/EEC. Enzootic Bovine Leukosis (EBL)

The agar gel immuno-diffusion test and the enzyme linked immuno-absorbent assay test (ELISA) shall be carried out according to paragraphs A and C of Chapter II of Annex D to Directive 64/432/EEC.

Bluetongue (BTG)

A. The blocking or competitive ELISA test shall be carried out according to the following protocol:

The competitive ELISA using monoclonal antibody 3-17-A3 is capable of identifying antibodies to all known serotypes of bluetongue virus (BTV).

The principle of the test is the interruption of the reaction between BTV antigen and a group-specific monoclonal antibody (3-17-A3) by the addition of test serum. Antibodies to BTV present in the test serum block the reactivity of the monoclonal antibody (Mab) and result in a reduction in the expected colour development after the addition of enzyme labelled anti-mouse antibody, and chromogen/substrate. Sera can be tested at a single dilution of 1:5 (spot test – Appendix 1) or may be titrated (serum titration – Appendix 2) to give dilution end-point. Inhibition values higher than 50 % may be regarded as positive.

Material and Reagents:

- 1. Appropriate ELISA microtitre plates.
- 2. Antigen: supplied as a cell extracted concentrate, prepared as described below, and stored at either -20 °C or -70 °C.
- 3. Blocking buffer: phosphate buffered saline (PBS) containing 0,3 % BTV negative adult bovine serum, 0,1 % (v/v) Tween-20 (supplied as polyoxyethylene sorbiton monolaurate syrup) in PBS.
- 4. Monoclonal antibody: 3-17-A3 (supplied as hybridoma tissue-culture supernatant) directed against the group-specific polypeptide VP7, stored at 20 °C or freeze-dried and diluted 1/100 with blocking buffer before use.
- 5. Conjugate: rabbit anti-mouse globulin (adsorbed and eluted) conjugated to horseradish peroxidase and kept in the dark at 4 °C.
- 6. Chromogen and substrate: Orthophenylene diamine (OPD-chromogen) at a final concentration of 0,4 mg/ml in sterile distilled water. Hydrogen peroxide (30 %w/v-substrate) 0,05 % v/v added immediately before use (5µl H₂ O₂ per 10 ml OPD). (Handle OPD with care wear rubber gloves suspected mutagen).
- 7. 1 Molar sulphuric acid: 26,6 ml of acid added to 473,4 ml of distilled water. (Remember Acid must be added to water, never water to acid.)
- 8. Orbital shaker.
- 9. ELISA plate reader (the test may be read visually).

Test format

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

Cc: conjugate control (no serum/ no monoclonal antibody); C++: strong positive serum control; C+: weak positive serum control; C-: negative serum control; Cm: monoclonal antibody control (no serum).

APPENDIX 1:

Spot dilution (1:5) format (40 sera/plate)

	Controls		Tes	t Sera								
	1	2	3	4	5	6	7	8	9	10	11	12
A	Cc	C-	1	2	3	4	5	6	7	8	9	10
В	Cc	C-	1	2	3	4	5	6	7	8	9	10
С	C++	C++										
D	C++	C++										
Е	C+	C+										
F	C+	C+										
G	Cm	Cm										40
Н	Cm	Cm										40

APPENDIX 2:

Serum titration format (10 sera/plate)

	Cont	Controls		Sera								
	1	2	3	4	5	6	7	8	9	10	11	12
A	Cc	C-	1:5									1:5
В	Cc	C-	1:10									1:10
С	C++	C++	1:20									1:20
D	C++	C++	1:40									1:40
Е	C+	C+	1:80									1:80
F	C+	C+	1:160									1:160
G	Cm	Cm	1:320									1:320
Н	Cm	Cm	1:640									1:640

Test protocol:

Conjugate control: Wells 1A and 1B are a blank control consisting of BTV antigen and

conjugate. This may be used to blank the ELISA reader.

Mab control (Cm)

Columns 1 and 2, rows G and H are the monoclonal antibody control and contain BTV antigen, monoclonal antibody and conjugate. These wells represent maximum colour. The mean of the optical density readings

from this control represents the 0 % inhibition value.

Positive control (C:

++, C+)

Columns 1 and 2, rows C-D-E-F. These wells contain BTV antigen, BTV strong and weak positive antiserum respectively, Mab and

conjugate.

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

Negative control: Wells 2A and 2B are the negative controls, which contain BTV antigen,

(C-)

BTV negative antiserum, Mab and conjugate.

Test sera : For large-scale serological surveys and rapid screening, sera may be

tested at a single dilution of 1:5 (Appendix 1). Alternatively, 10 sera may be tested over a dilution range from 1:5 to 1:640 (Appendix 2). This will give some indication of the titre of antibody in the test sera.

Procedure:

1. Dilute BTV antigen to pre-titrated concentration in PBS, sonicate briefly to disperse aggregated virus (if sonicator is not available, pipette vigorously) and add 50 μ l to all wells of the ELISA plate. Tap sides of plate to disperse antigen.

- 2. Incubate at 37 °C for 60 minutes on an orbital shaker. Wash plates three times by flooding and emptying the wells with non-sterile PBS and blot dry on absorbent paper.
- 3. Control wells: Add 100 μl of blocking buffer to Cc wells. Add 50 ul of positive and negative control sera, at a dilution of 1:5 (10 μ l sera + 40 μl blocking buffer), to respective wells C-, C+ and C++. Add 50μl blocking buffer to Mab control wells.

Spot titration method: Add a 1:5 dilution of each test serum in blocking buffer to duplicate wells of columns 3 to 12 (10 µl sera + 40 µl blocking buffer),

or

Serum titration method: Prepare a two-fold dilution series of each test sample (1:5 to 1:640) in blocking buffer across eight wells of single columns 3 to 12.

- 4. Immediately after the addition of the test sera, dilute Mab 1:100 in blocking buffer and add 50 μl to all wells of the plate except for the blank control.
- 5. Incubate at 37 °C for 60 minutes on an orbital shaker. Wash three times with PBS and blot dry.
- 6. Dilute rabbit anti-mouse concentrate to 1/5~000 in blocking buffer and add $50~\mu l$ to all wells of the plate.
- 7. Incubate at 37 °C for 60 minutes on an orbital shaker. Wash three times with PBS and blot dry.
- 8. Thaw the O-Phenylenediamine dihydrochloride (OPD) and immediately before use add 5 µl of 30 % hydrogen peroxide to each 10 ml of OPD. Add 50 µl to all wells of the plate. Allow colour to develop for approximately 10 minutes and stop the reaction with 1 Molar sulphuric acid (50 µl per well). Colour should develop in the Mab control wells and in those wells containing sera with no antibody to BTV.
- 9. Examine and record the plates either visually or using a spectrophotometric reader. Analysis of results:

Using the software package print out the optical density (OD) values, and the percentage inhibition (PI) for test and control sera based on the mean value recorded in the antigen control wells. The date expressed as OD and PI values are used to determine whether the test has performed within acceptable limits. The upper control limits (UCL) and lower control limits (LCL) for the Mab control (antigen plus Mab in the absence of test sera) are between OD values 0,4 and 1.4. Any plate that fails to conform to the above criteria must be rejected.

If a computer software package is not available print out the OD values using the ELISA printer. Calculate the mean OD value for the antigen control wells, which is equivalent to the 100 %

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

value. Determine the 50 % OD value and manually calculate the positivity and negativity of each sample.

Percentage inhibition (PI) value = $100 - (OD \text{ of each test control/Mean OD of Cm}) \times 100$.

The duplicate negative control serum wells and the duplicate blank wells must record PI values between +25% and -25%, and between +95% and +105%, respectively. Failure to be within these limits does not invalidate the plate but does suggest that background colour is developing. The strong and weak positive control sera must record PI values between +81% and +100%, and between +51% and +80%, respectively.

The diagnostic threshold for test sera is 50 % (PI 50 % or OD 50 %). Samples recording PI values >50 % are recorded negative. Samples that record PI values above and below the threshold for the duplicate wells are considered doubtful; such samples may be re-tested in the spot test and/or titration. Positive samples may also be titrated to provide an indication of the degree of positivity.

Visual reading: Positive and negative samples are easily discernible by eye; weakly positive or strong negative samples may be more difficult to interpret by eye.

Preparation of BTV ELISA antigen:

- 1. Wash 40-60 roux of confluent BHK-21 cells three times with serum-free Eagle's medium and infect with bluetongue virus serotype 1 in serum-free Eagle's medium.
- 2. Incubate at 37 °C and examine daily for cytopathic effect (CPE).
- 3. When CPE are complete in 90 % to 100 % of the cell sheet of each roux, harvest the virus by shaking any still-attached cells from the glass.
- 4. Centrifuge at 2 000 to 3 000 rpm to pellet the cells.
- 5. Discard the supernatant and re-suspend the cells in approximately 30 ml of PBS containing 1 % 'Sarkosyl' and 2 ml phenylmethylsulphonyl fluoride (lysis buffer). This may cause the cells to form a gel and more lysis buffer may be added to reduce this effect. (NB: phenylmethylsulphonyl fluoride is harmful handle with extreme caution.)
- 6. Disrupt the cells for 60 seconds using an ultrasonic probe at an amplitude of 30 microns.
- 7. Centrifuge at 10 000 rpm for 10 minutes.
- 8. Store the supernatant at + 4 °C and re-suspend the remaining cell pellet in 10 to 20 ml of lysis buffer.
- 9. Sonicate and clarify, storing the supernatant at each stage, a total of three times.
- 10. Pool the supernatants and centrifuge at 24 000 rpm (100,000 g) for 120 minutes at + 4 °C over a 5 ml cushion of 40 % sucrose (w/v in PBS) using 30 ml Beckmann centrifuge tubes and an SW 28 rotor.
- Discard the supernatant, drain the tubes thoroughly and re-suspend the pellet in PBS by sonication. Store the antigen in aliquots at -20 °C.

Titration of BTV ELISA antigen:

Bluetongue ELISA antigen is titrated by the indirect ELISA. Twofold dilutions of antigen are titrated against a constant dilution (1/100) monoclonal antibody 3-17-A3. The protocol is as follows:

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

- 1. Titrate a 1:20 dilution of BTV antigen in PBS across the microtitre plate in a twofold dilution series (50 µl/well) using a multichannel pipette.
- 2. Incubate for one hour at 37 °C on an orbital shaker.
- 3. Wash plates three times with PBS.
- 4. Add 50 μ l of monoclonal antibody 3-17-A3 (diluted 1/100) to each well of the microtitre plate.
- 5. Incubate for one hour at 37 °C on an orbital shaker.
- 6. Wash plates three times with PBS.
- 7. Add 50 μl of rabbit anti-mouse globulin conjugated to horseradish peroxidase, diluted to a pre-titrated optimal concentration, to each well of the microtitre plate.
- 8. Incubate for one hour at 37 °C on an orbital shaker.
- 9. Add substrate and chromogen as described previously. Stop the reaction after 10 minutes by the addition of 1 Molar sulphuric acid (50 μl/well).

In the competitive assay, the monoclonal antibody must be in excess, therefore a dilution of antigen is chosen which falls on the titration curve (not on the plateau region) which gives approximately 0,8 OD after 10 minutes.

B. The agar gel immuno-diffusion test shall be carried out according to the following protocol:

Antigen:

Precipitating antigen is prepared in any cell culture system that supports the rapid multiplication of a reference strain of bluetongue virus. BHK or Vero cells are recommended. Antigen is present in the supernatant fluid at the end of virus growth but requires 50 to 100-fold concentration to be effective. This may be achieved by any standard protein concentration procedure; virus in the antigen may be inactivated by the addition of 0.3% (v/v) beta-propiolactone.

Known positive control serum:

Using the international reference serum and antigen a national standard serum is produced, standardised for optimal proportion against the international reference serum, freeze-dried and used as the known control serum in each test.

Test serum

Procedure : 1 % agarose prepared in borate or sodium barbitol buffer, pH 8,5 to 9,0,

is poured into a petri dish to a minimum depth of 3,0 mm. A test pattern of seven moisture-free wells, each 5,0 mm in diameter, is cut in the agar. The pattern consists of one centre well and six wells arranged round it in a circle of radius 3 cm. The central well is filled with the standard antigen. Peripheral wells 2, 4 and 6 are filled with known positive serum, wells 1, 3 and 5 are filled with test sera. The system is incubated for up

to 72 hours at room temperature in a closed humid chamber.

Interpretation : A test serum is positive if it forms a specific precipitin line with the

antigen and forms a complete line of identity with the control serum. A test serum is negative if it does not form a specific line with the antigen and it does not bend the line of the control serum. Petri dishes must be examined against a dark background and using indirect illumination.

Epizootic haemorrhagic disease (EHD)

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

The agar gel immuno-diffusion test shall be carried out according to the following protocol: Antigen:

Precipitating antigen is prepared in any cell culture system that supports the rapid multiplication of the appropriate serotype(s) of epizootic haemorrhagic disease virus. BHK or Vero cells are recommended. Antigen is present in the supernatant fluid at the end of virus growth but requires 50 to 100-fold concentration to be effective. This may be achieved by any standard protein concentration procedure; virus in the antigen may be inactivated by the addition of 0,3% (v/v) beta-propiolactone.

Known positive control serum:

Using the international reference serum and antigen a national standard serum is produced, standardised for optimal proportion against the international reference serum, freeze-dried and used as the known control serum in each test.

Test serum

Procedure : 1 % agarose prepared in borate or sodium barbitol buffer, pH 8,5 to 9,0,

is poured into a petri dish to a minimum depth of 3,0 mm. A test pattern of seven moisture-free wells, each 5,0 mm in diameter, is cut in the agar. The pattern consists of one centre well and six wells arranged round it in a circle of radius 3 cm. The central well is filled with the standard antigen. Peripheral wells 2, 4 and 6 are filled with known positive serum, wells 1, 3 and 5 are filled with test sera. The system is incubated for up

to 72 hours at room temperature in a closed humid chamber.

Interpretation : A test serum is positive if it forms a specific precipitin line with the

antigen and forms a complete line of identity with the control serum. A test serum is negative if it does not form a specific line with the antigen and it does not bend the line of the control serum. Petri dishes must be examined against a dark background and using indirect illumination.

Infectious bovine rhinotracheitis (IBR) / infectious pustular vulvo-vaginitis (IPV)

A. The serum neutralisation test shall be carried out according to the following protocol:

Serum : All sera are heat-inactivated at 56 °C for 30 minutes before use.

Procedure : The constant virus-varying serum neutralisation test on microtitre plates

employs MDBK or other susceptible cells. The Colorado, Oxford or any other reference strain of the virus is used at 100 TCID50 per 0,025 ml; inactivated undiluted serum samples are mixed with an equal volume (0,025 ml) of virus suspension. The virus/serum mixtures are incubated for 24 hours at 37 °C in the microtitre plates before the MDBK cells are added. Cells are used at a concentration which forms a complete

monolayer after 24 hours.

Controls : (i) virus infectivity assay, (ii) serum toxicity controls, (iii) uninoculated

cell culture controls, (iv) reference antisera.

Interpretation : The results of the neutralisation test and the titre of the virus used in the

test are recorded after three to six days incubation at 37 $^{\circ}$ C. Serum titres are considered negative if there is no neutralisation at a dilution of 1/2

(undiluted serum).

B. Any other test recognised in the framework of Decision 2004/558/EC⁽⁵⁾. Foot-and-mouth disease (FMD)

A. Collecting oesophageal/pharyngeal samples and testing shall be carried out according to the following protocol:

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

Reagents

Prior to sampling, transport medium is prepared. Two ml volumes are dispensed in as many containers as there are animals to be sampled. The containers used must withstand freezing over solid CO₂ or liquid nitrogen. Samples are obtained by the use of a specially-designed sputum collector or 'probang'. To obtain a sample the probang cup is passed through the mouth, over the dorsum of the tongue and down into the upper part of the oesophagus. Attempts are made to scrape the surface epithelium of the upper oesophagus and pharynx by movements directed laterally and dorsally. The probang is then withdrawn, if possible after the animal has swallowed. The cup must be full and contain a mixture of mucus, saliva, oesophageal fluid and cellular debris. Care must be taken to ensure that each specimen contains some visible cellular material. Very rough handling which causes bleeding must be avoided. Samples from some animals may be heavily contaminated with ruminal contents. Such samples must be discarded and the mouth of the animal flushed with water, or preferably physiological saline, before repeat sampling.

Treatmentof samples:

Each sample collected in the probang cup is examined for quality and 2 ml added to an equal volume of transport medium in a container which can withstand freezing. The containers are tightly closed, sealed, disinfected and labelled. The samples are kept cool (+ 4 °C) and examined within three to four hours or placed over dry ice (- 69 °C) or liquid nitrogen and kept frozen until examined. Between animals the probang is disinfected and washed in three changes of clean water.

Testing for FMD : virus:

Samples are inoculated into cultures of primary bovine thyroid cell cultures using at least three tubes per sample. Other susceptible cells such as primary bovine or porcine kidney cells can be used but it must be kept in mind that for some strains of FMD virus they are less sensitive. The tubes are incubated at 37 °C on a roller apparatus and examined daily for 48 hours for the presence of a cytopathic effect (CPE). If negative, cultures are blind passaged onto new cultures and re-examined for 48 hours. The specificity of any CPE must be confirmed.

Recommended transport media:

- 1. 0,08M phosphate buffer pH 7,2 containing 0,01 % bovine serum albumin, 0,002 % phenol red and antibiotics.
- 2. Tissue culture medium (such as Eagle's MEM) containing 0,04 M Hepes buffer, 0,01 % bovine serum albumin and antibiotics, pH 7,2.
- 3. Antibiotics (per ml final) must be added to the transport medium such as penicillin 1 000 IU, neomycin sulphate100 IU, polymyxin B sulphate50 IU, mycostatin100 IU.
- B. The virus neutralisation test shall be carried out according to the following protocol:

Reagents

Stock FMDV antigen is prepared in cell cultures or on cattle tongues and stored at - 70 °C or less or at - 20 °C after the addition of 50 % glycerol. This is the stock antigen. FMDV is stable under these conditions and titres vary little over a period of months.

Procedure

The test is carried out in flat-bottomed tissue culture grade microtitre plates using susceptible cells such as IB-RS-2, BHK-21 or calf kidney cells. Sera for the test are diluted 1/4 in serum-free cell culture medium with the addition of 100 IU/ml neomycin or other suitable antibiotics. Sera are inactivated at 56 °C for 30 minutes and 0.05 ml amounts

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

are used to prepare a twofold series on microtitre plates using 0,05 ml diluting loops. Pre-titrated virus also diluted in serum-free culture medium and containing 100 TCID50/0.05 ml is then added to each well. Following incubation at 37 °C for one hour to allow neutralisation to take place, 0,05 ml of suspension cells containing 0,5 to 1.0×10^6 cells per 1 ml in cell culture medium containing serum free of FMD antibody is added to each well and the plates are sealed. Plates are incubated at 37 °C. Monolayers are normally confluent within 24 hours. CPE is usually sufficiently advanced at 48 hours for a microscopic reading of the test. At this time a final microscopic reading may be made or the plates may be fixed and stained for macroscopic reading, for instance using 10 % formol-saline and 0,05 % methylene blue.

Controls

Controls in each test include homologous antiserum of known titre, a cell control, a serum toxicity control, a medium control, and a virus titration from which the actual amount of virus in the test is calculated.

Interpretation

Wells with evidence of CPE are considered to be infected and neutralisation titres are expressed as the reciprocal of the final dilution of serum present in the serum/virus mixtures at the 50 % end point estimated according to the Spearman-Karber method. (Karber, G., 1931, Archiv fuer Experimentelle Pathologie und Pharmokologie, 162, 480.). Tests are considered to be valid when the actual amount of virus used per well in the test is between 101,5 and 102,5 TCID50 and when the titre of the reference serum is within twofold of its expected titre, estimated from the mode of previous titrations. When the controls are outside these limits the tests are repeated. An end point titre of 1/11 or less is taken as negative.

C. The detection and quantification of antibody by ELISA shall be carried out according to the following protocol:

Reagents

: Rabbit antisera to 146S antigen of seven types of foot-and-mouth disease virus (FMDV) used at a predetermined optimum concentration in carbonate/bicarbonate buffer, pH 9,6. Antigens are prepared from selected strains of virus grown on monolayers of BHK-21 cells. The unpurified supernatants are used and pretitrated according to the protocol but without serum, to give a dilution which after the addition of an equal volume of PBST (phosphate buffered saline containing 0,05 % Tween-20 and phenol red indicator) would give an optical density reading of between 1,2 and 1,5. The viruses can be used inactivated. PBST is used as a diluent. Guinea-pig antisera are prepared by inoculating guinea pigs with 146S antigen of each serotype. A predetermined optimum concentration is prepared in PBST containing 10 % normal bovine serum and 5 % normal rabbit serum. Rabbit antiguinea-pig immunoglobulin conjugated to horseradish peroxidase is used at a predetermined optimum concentration in PBST containing 10 % normal bovine serum and 5 % normal rabbit serum. Test sera are diluted in PBST.

Procedure:

- 1. ELISA plates are coated with 50 μ l of rabbit antiviral sera overnight in a humidity chamber at room temperature.
- 2. Fifty microlitres of a duplicate, twofold series of each test serum starting at 1/4 are prepared in U-bottomed multiwell plates (carrier plates). Fifty microlitres of a constant

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

dose of antigen are added to each well and the mixtures are left overnight at 4 °C. The addition of the antigen reduces the starting serum dilution to 1/8.

- 3. The ELISA plates are washed five times with PBST.
- 4. Fifty microlitres of serum/antigen mixtures are then transferred from the carrier plates to the rabbit-serum-coated ELISA plates and incubated at 37 °C for one hour on a rotary shaker.
- 5. After washing, 50 μl of guinea-pig antiserum to the antigen used in point 4 is added to each well. The plates are incubated at 37 °C for one hour a rotary shaker.
- 6. The plates are washed and 50 μl of rabbit anti-guinea-pig immunoglobulin conjugated to horseradish peroxidase is added to each well. The plates are incubated at 37 °C for one hour on a rotary shaker.
- 7. The plates are washed and 50 μ l of orthophenylene diamine containing 0,05 % H_2O_2 (30 %) w/v is added to each well.
- 8. The reaction is stopped after 15 minutes with 1,25M H₂SO₄.

The plates are read spectrophotometrically at 492 nm on an ELISA reader linked to a microcomputer.

Controls : For each antigen used 40 wells contain no serum but contain antigen

diluted in PBST. A duplicated twofold dilution series of homologous bovine reference antiserum. A duplicate twofold dilution series of

negative bovine serum.

Interpretation : Antibody titres are expressed as the final dilution of tests serum giving

50 % of the mean OD value recorded in the virus control wells where test serum is absent. Titres in excess of 1/40 are considered positive.

References : Hamblin C, Barnett ITR and Hedger RS (1986) 'A new enzyme-linked

immunosorbent assay (ELISA) for the detection of antibodies against foot-and-mouth disease virus. I. Development and method of ELISA.'

Journal of Immunological Methods, 93, 115 to 121.11.

Aujeszky's disease (AJD)

A. The serum neutralisation test shall be carried out according to the following protocol:

Serum : All sera are heat-inactivated at 56 °C for 30 minutes before use.

Procedure : The constant virus-varying serum neutralisation test on microtitre plates

employs Vero or other sensitive cell systems. Aujeszky's disease virus is used at 100 TCID50 per 0,025 ml; inactivated undiluted serum samples are mixed with an equal volume (0,025 ml) of virus suspension. The virus/serum mixtures are incubated for two hours at 37 °C in the microtitre plates before the appropriate cells are added. Cells are used at a concentration which forms a complete monolayer after 24 hours.

Controls : (i) virus infectivity assay, (ii) serum toxicity controls, (iii) uninoculated

cell culture controls, (iv) reference antisera.

Interpretation : The results of the neutralisation test and the titre of the virus used in the

test are recorded after three to seven days incubation at 37 °C. Serum

titres less than 1/2 (undiluted serum) are considered negative.

B. Any other test recognised in the framework of Decision 2008/185/EC⁽⁶⁾. Transmissible gastro-enteritis (TGE)

The serum neutralisation test shall be carried out according to the following protocol:

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

Serum : All sera are heat-inactivated at 56 °C for 30 minutes before use.

Procedure : The constant virus-varying serum neutralisation test on microtitre plates

employs A72 (dog tumour) cells or other sensitive cell systems. TGE virus is used at 100 TCID50 per 0,025 ml; inactivated undiluted serum samples are mixed with an equal volume (0,025 ml) of virus suspension. The virus/serum mixtures are incubated for 30 to 60 minutes at 37 °C in the microtitre plates before the appropriate cells are added. Cells are used at a concentration which forms a complete monolayer after 24

hours. Each cell receives 0,1 ml of cell suspension.

Controls : (i) virus infectivity assay, (ii) serum toxicity controls, (iii) uninoculated

cell culture controls, (iv) reference antisera.

Interpretation : The results of the neutralisation test and the titre of the virus used in

the test are recorded after three to five days incubation at $37\,^{\circ}$ C. Serum titres less than 1/2 (final dilution) are considered negative. If undiluted serum samples are toxic to the tissue cultures, these sera may be diluted 1/2 before being used in the test. This is equivalent to 1/4 final dilution of serum. Serum titres of less than 1/4 (final dilution) are considered

negative in these cases.

Swine vesicular disease (SVD)

Tests for swine vesicular disease (SVD) shall be carried out according to Decision 2000/428/ EC⁽⁷⁾.

Classical swine fever (CSF)

Tests for classical swine fever (CSF) shall be carried out according to Decision 2002/106/EC⁽⁸⁾.

The performance of tests for CSF must follow the guidelines set out in the relevant chapter of the OIE Manual of Diagnostic Tests and Vaccines for Terrestrial Animals.

The evaluation of sensitivity and specificity of the serological test for CSF must be carried out in a national laboratory with a quality assurance scheme in place. Tests employed must be shown to recognise a range of weak and strong positive reference sera and allow detection of antibody in early phase and convalescence.

I^{F5}Vesicular stomatitis (VS)

The virus neutralisation (VN) test shall be carried out in accordance with the testing protocols for vesicular stomatitis set out in Chapter 2.1.19 of the OIE Manual of Diagnostic Tests and Vaccines for Terrestrial Animals.

Sera that prevent cytopathic effect (CPE) at dilutions of 1 in 32 or greater shall be considered to contain antibodies to the vesicular stomatitis virus.]

PART 7

Import and quarantine animal health conditions for animals imported into St. Pierre and Miquelon within a period of less than six months prior to introduction into the Union

(referred to in Article 6)

Animal species covered

Taxon		
ORDER	FAMILY	GENUS AND SPECIES

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

Artiodactyla	Camelidae	Camelus spp., Lama spp.,
Titiodactyla		Vicugna spp.,

CHAPTER 1

Residence and quarantine

- 1. Animals imported into St. Pierre and Miquelon must reside in an authorised quarantine station for a minimum period of 60 days before being dispatched for introduction into the Union. This period may be increased due to testing requirements for individual species. In addition the animals must comply with the following requirements:
- (a) Separate consignments may enter the quarantine station. However, upon entry in the quarantine station all animals of the same species in the quarantine facility must be considered as a single group, and referred to as such. The quarantine period must commence for the whole group at the time when the last animal entered the quarantine facility.
- (b) Within the quarantine station each specific group of animals must be maintained in isolation, with no direct or indirect contact with any other animals, including those from other consignments that may be present.
 - Each consignment must be kept in the approved quarantine station and protected from vector insects.
- (c) If, during the period of quarantine, the isolation of a group of animals is not maintained and contact is made with other animals, the quarantine period must begin again for the same duration as initially prescribed on entry into the quarantine station.
- (d) Animals to be introduced into the Union which pass through the quarantine station must be loaded and dispatched directly to the Union:
 - (i) without coming into contact with animals other than animals which fulfil the health conditions established for the introduction of the relevant category of animal into the Union;
 - segregated into consignments so that no consignment can came in contact with animals not eligible for importation into the Union;
 - (iii) in transport vehicles or containers which have first been cleansed and disinfected with a disinfectant officially authorised in St. Pierre and Miquelon as effective in the control of the diseases referred to in Chapter 2 and which are so constructed that faeces, urine, litter or fodder cannot flow or fall out of the vehicle or container during transportation.
- 2. The quarantine premises must at least meet the minimum standards laid down in Annex B to Directive 91/496/EEC⁽⁹⁾, and the following conditions:
- (a) they must be supervised by an official veterinarian;
- (b) they must be situated at the centre of an area of at least 20 km in diameter in which, according to official findings, for at least 30 days prior to their use as a quarantine station there has been no case of foot-and-mouth disease;

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

- (c) they must, before being used as a quarantine station, be cleansed and disinfected with a disinfectant officially authorised in St Pierre et Miquelon as effective in the control of the diseases referred to in Chapter 2;
- (d) they must operate, taking into account their animal capacity:
 - (i) a facility dedicated exclusively for the quarantine of animals, including adequate housing to a suitable standard for the animals;
 - (ii) appropriate facilities, that:
 - are easy to thouroughly clean and disinfect,
 - include facilities for safe loading and unloading,
 - are able to fulfil all watering and feeding requirements for the animals,
 - allow any necessary veterinary treatment to be easily administered;
 - (iii) appropriate facilities for inspection and isolation;
 - (iv) appropriate equipment for cleaning and disinfecting rooms and transport vehicles:
 - (v) an appropriate storage area for fodder, litter and manure;
 - (vi) an appropriate system for collecting waste water;
 - (vii) an office for the official veterinarian;
- (e) when operating, they must have sufficient veterinarians to carry out all duties;
- they must only admit animals that are individually identified so as to guarantee traceability. To this end, when animals are admitted the owner or the person in charge of the quarantine station must ensure that the animals are properly identified and accompanied by health certificates for the species and categories involved. In addition, the owner or the person in charge of the quarantine station must record on a register or in a data base, and retain for at least three years, the name of the owner, the origin of the animals in the consignment, the identification number of the animals in the consignment and their place of destination;
- (g) the competent authority must determine the procedure for official supervision of the quarantine station and must ensure that such supervision is carried out; this supervision must include regular inspections in order to ascertain that the requirements for approval continue to be fulfilled. In case of failure and suspension, the approval may only be restored when the competent authority is satisfied that the quarantine premises are in full compliance with all the conditions set out in points (a) to (g).

CHAPTER 2

Animal health tests

1. GENERAL REQUIREMENTS

The animals must be subjected to the following tests carried out on samples of blood taken, if not specified otherwise, not earlier than 21 days from the date of commencement of the isolation period.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

The laboratory tests must be carried out in an approved laboratory in the Union and all laboratory tests and their results, vaccinations and treatments must be enclosed with the health certificate.

In order to keep animal interventions to a minimum, sampling, tests and any vaccinations must be grouped as far as is possible whilst respecting the minimum time intervals required by the testing protocols set out in Part 2 of this Chapter.

2. SPECIFIC REQUIREMENTS

2.1 CAMELIDAE

2.1.1 Tuberculosis

(a) **Test to be used**: comparative intradermal reaction test using Bovine purified protein derivative (PPD) and Avian PPD conforming to the standards for the manufacture of bovine and avian tuberculins as described in point 2.1.2 of Annex B of Directive 64/432/EEC.

The test must be executed in the area behind the shoulder (axillary region) following the technique described in point 2.2.4 of Annex B of Directive 64/432/EEC.

(b) **Timing**: the animals must be tested within two days from the date of arrival in the quarantine station and 42 days from the date of the first test.

(c) Interpretation of tests:

the reaction shall be considered:

- negative if the increased skin thickness is less than 2 mm.
- positive if the increased skin thickness is more than 4 mm.
- inconclusive if the increased skin thickness to the bovine PPD is between 2mm and 4 mm, or more than 4 mm but less then the reaction to the avian PPD.

(d) Options for action following testing:

If an animal presents a positive result to the intradermal-reaction to the bovine PPD, that animal shall be excluded from the group and the other animals shall be re-tested starting at least 42 days from the date of the first positive test was administered and this shall be considered as the first test described in (b).

If more than one animal of the group presents a positive result, the whole group shall be rejected for exportation to the Union.

If one or more animals of the same group present an inconclusive reaction, the whole group shall be re-tested starting at least 42 days from the date of the first test was administered and it shall be considered as the first test described in (b).

2.1.2 Brucellosis

(a) Test to be used:

(i) Brucella abortus: Rose Bengal test (RBT) and Serum agglutination test (SAT) as described respectively in points 2.5 and 2.6 of Annex C to Directive 64/432/EEC. In the case of a positive result, a complement-fixation test shall be performed for confirmation as described in Part 6 of Annex I to Regulation (EU) No 206/2010.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

- (ii) Brucella melitensis: RBT and SAT as described respectively in points 2.5 and 2.6 of Annex C to Directive 64/432/EEC. In the case of a positive result, a complement-fixation test following the method described in Annex C to Directive 91/68/EEC shall be performed for confirmation.
- (iii) Brucella ovis: Complement fixation test as described in Annex D to Directive 91/68/EEC
- (b) **Timing**: the animals have to be tested within two days from the date of their arrival in the quarantine station and 42 days from the date of the first test.
- (c) Interpretation of tests:

A positive reaction to the tests shall be as defined in Annex C to Directive 64/432/EEC.

(d) Options for action following testing:

Animals tested positive to one of the tests shall be excluded from the group and the other animals shall be re-tested starting at least 42 days from the date the first positive test was performed: this shall be considered as the first test described in (b).

Only the animals that tested negative to two consecutive tests performed as described in (b) shall be allowed for the introduction to the Union.

- 2.1.3 Bluetongue and Epizootic haemorrhagic disease (EHD)
- (a) **Test to be** used: agar gel immunodiffusion (AGID) test as described in Part 6 of Annex I to Regulation (EU) No 206/2010.

In case of a positive reaction the animals shall be tested with competitive ELISA test as described in Part 6 of Annex I to Regulation (EU) No 206/2010 to discriminate between the two diseases.

(b) Timing:

The animals must be tested with negative result to two tests: the first within two days from the date of their arrival in the quarantine station and the second at least 21 days from date of the first test.

(c) Options for action following testing:

(i) Bluetongue

If one or more animals tested positive to the ELISA as described in Part 6 of Annex I to Regulation (EU) No 206/2010, the positive animal/animals shall be excluded from the group, and all the remaining animals in the group must be quarantined for 100 days starting from the date on which the samples for the positive test were collected. The group shall only be considered free of the bluetongue disease if regular checks carried out by official veterinarians throughout the duration of the quarantine period fail to reveal clinical symptoms of disease, and the quarantine station remains free of bluetongue vectors (*Culicoides*).

If a further animal presents clinical symptoms of bluetongue disease during the quarantine period as described in the first subparagraph, all the animals in the group shall be rejected for introduction into the Union.

(ii) Epizootic haemorrhagic disease (EHD)

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

If one or more animals tested positive reveal the presence of antibodies to the EHD virus during confirmatory ELISA testing, the animal(s) shall be considered positive and shall be excluded from the group, and the whole group shall be subject to repeat testing beginning at least 21 days from the date of the initial positive diagnosis and again at least 21 days from the date of the repeat test, both with negative results.

If any additional animals are tested positive during either or both of the two tests carried out for repeat testing, the whole group of animals shall be rejected for introduction into the Union.

2.1.4 Foot-and-Mouth Disease (FMD)

- (a) **Test to be used**: Diagnostic tests (probang and serology) using ELISA and (Virus Neutralisation) (VN) techniques in accordance with the Protocols described in Part 6 of Annex I to Regulation (EU) No 206/2010.
- (b) **Timing**: the animals shall be tested with negative results to two tests: the first within two days from the date of their arrival in the quarantine station and the second at least 42 days from the date of the first test.
- (c) **Options for action following testing**: If any animal tests positive for the FMD virus, then none of the animals present in the quarantine station shall be considered as eligible for introduction into the Union.

Note: Any detection of antibodies to structural or not structural proteins of FMD virus shall be considered as a result of previous infection of FMD irrespective of the vaccination status.

2.1.5 Rinderpest

- (a) **Test to be used**: The competitive ELISA test as described in the OIE Manual of Diagnostic Tests and Vaccines for Terrestrial Animals, latest version, is the prescribed test for international trade and is test of choice. Serum neutralisation test, or other recognised tests in accordance with the protocols described in relevant sections of the OIE manual may also be used.
- (b) **Timing**: the animals shall be tested twice: the first within two days from the date of their arrival in the quarantine station and the second at least 42 days from the date of the first test.
- (c) **Options for action following testing**: If any animal tests positive for the Rinderpest virus, then none of the animals present in the quarantine station shall be considered as eligible for introduction into the Union.

2.1.6 Vesicular stomatitis

- (a) **Test to be used**: ELISA, virus neutralisation test, or other recognised test in accordance with the protocols described in the relevant sections of the OIE manual.
- (b) **Timing**: the animals shall be tested twice: the first within two days from the date of their arrival in the quarantine station and the second at least 42 days from the date of the first test.
- (c) **Options for action following testing**: If any animal tests positive for vesicular stomatitis virus, then none of the animals present in the quarantine station shall be considered as eligible for introduction into the Union.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

- 2.1.7 Rift valley fever
- (a) **Test to be used**: ELISA, virus neutralisation test, or other recognised test in accordance with the protocols described in relevant sections of the OIE manual.
- (b) **Timing**: the animals shall be tested twice: the first within two days from the date of their arrival in the quarantine station and the second at least 42 days from the date of the first test.
- (c) **Options for action following testing**: If any animal displays evidence of exposure to rift valley fever agent, then none of the animals present in the quarantine station shall be considered as eligible for introduction into the Union.
- 2.1.8 Lumpy skin disease
- (a) **Test to be used**: Serology using ELISA, virus neutralisation test, or other recognised test in accordance with the protocols described in relevant sections of the OIE manual.
- (b) **Timing**: the animals shall be tested twice: the first within two days from the date of their arrival in the quarantine station and the second at least 42 days from the date of the first test.
- (c) **Options for action following testing**: If any animal displays evidence of exposure to lumpy skin disease, then none of the animals present in the quarantine station shall be considered as eligible for introduction into the Union.
- 2.1.9 Crimean congo haemorrhagic fever
- (a) **Test to be used**: ELISA, virus neutralisation test, Immunofluorescence test or other recognised test.
- (b) **Timing**: the animals shall be tested twice: the first within two days from the date of their arrival in the quarantine station and the second at least 42 days from the date of the first test.
- (c) **Options for action following testing**: If any animal displays evidence of exposure to crimean congo haemorrhagic fever agent, then none of the animals present in the quarantine station shall be considered as eligible for introduction into the Union.
- 2.1.10 Surra (Trypanosoma evansi (T. evansi))
- (a) **Test to be used**: The parasitic agent can be identified in concentrated blood samples in accordance with the protocols described in relevant sections of the OIE manual.
- (b) **Timing**: the animals shall be tested twice: the first within two days from the date of their arrival in the quarantine station and the second at least 42 days from the date of the first test.
- (c) **Options for action following testing**: If *T. evansi* is detected in any animal in the consignment, then that animal shall be considered not eligible for introduction into the Union. The remaining animals of the group shall then undergo internal and external antiparasitic treatment using suitable agents that are effective against *T. evansi*.
- 2.1.11 Malignant catarrhal fever
- (a) **Test to be used**: Detection of viral DNA based on identification by immunofluorescence or immunocytochemistry using the protocols described in relevant sections of the OIE manual.

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

- (b) **Timing**: the animals shall be tested twice: the first within two days from the date of their arrival in the quarantine station and the second at least 42 days from the date of the first test.
- (c) **Options for action following testing**: If any animal displays evidence of exposure to MCF, then none of the animals present in the quarantine station shall be considered as eligible for introduction into the Union.

2.1.12 Rabies

Vaccination: Rabies vaccination may be carried out when requested by the Member State of destination and the animal shall be blood sampled and a serum neutralisation test for antibodies carried out.

- 2.1.13 *Enzootic bovine leucosis*. (only in the case where the animals are destined for an officially enzootic-bovine-leucosis free Member State or region, as referred to in Article 2(2)(k) of Directive 64/432/EEC)
- (a) **Test to be used**: AGID or blocking ELISA, in accordance with the protocols described in the OIE manual, latest version.
- (b) **Timing**: the animals shall be tested twice: the first within two days from the date of their arrival in the quarantine station and the second at least 42 days from the date of the first test.
- (c) **Options for action following testing**: animals tested positive to the test described in (a) shall be excluded from the group of animals in the quarantine facility and the other animals shall be re-tested starting at least 21 days from the date of the first positive test was performed: this shall be considered as the first test described in (b).

Only the animals that tested negative to two consecutive tests performed as described in (b) shall be considered eligible for introduction into the Union.

ANNEX II

FRESH MEAT

[F12PART 1

LIST OF THIRD COUNTRIES, TERRITORIES AND PARTS THEREOF⁰

ISO Code of Code of Territo			onVeterinary certificate		Specific condition	Closing s date ^b	Opening date ^c	
and name of third country		country, territory or part thereof	Model(s)	SG				
1	2	3	4	5	6	7	8	
AL – Albania	AL-0	Whole country	_					

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

[F15AR —	AR-0	Whole	EQU				
Argentina		country					
\mathcal{L}	AR-1	The	BOV	A	1		1 Angust
	AK-I		RUF	A	1		1 August 2010
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			ormosa,				
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		te	rritory				
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	ir A	R-3).		
AR-2	The provinces of: CS SC T dd dF P or SC T dC T T T T T T T T T T T T T T T T T	BOV OVI RUW IRIbut, anta ruz, ierra el uego, art f feuquén except onfluencia ae one ocated ast f ie rovincial oad 7, ad icun eufú ae one ocated ast f ie rovincial oad 77, ad icun eufú ae one ocated ast f ie rovincial oad 77, ad icun eufú ae one ocated ast f ie rovincial oad 77, ad icun eufú ae one ocated ast f ie rovincial oad 77,		1 August 2008
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Department			
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San			
Antonio			
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Status: Point in time view as at 01/07/2016.

		zo lo es o th P	ne rovincial pads 50 nd			
	AR-3	Part of Salta: the area of 25 km from the border with Bolivia and Paraguay that extends from the Santa Catalina District in the Province of Jujuy, to the Laishi District in the Province of Formosa (the former high- surveillanc buffer area)	BOV RUF RUW	A		1 July 2016]
AU – Australia	AU-0	Whole country	BOV, OVI, POR, EQU, RUF, RUW, SUF, SUW			

BA – Bosnia and Herzegovii	BA-0	Whole country				
BH – Bahrain	BH-0	Whole country	_			
[^{F15} BR — BRAZIL	BR-0	Whole country	EQU			
	BR-1	State of Minas Gerais, State of Espírito Santo, State of Goiás, State of Mato Grosso, State of Rio Grande Do Sul, State of Mato Grosso Do Sul (excluding territory included in BR-4).	BOV	A and H	1	1 December 2008
	BR-2	State of Santa Catarina	BOV	A and H	1	31 January 2008
	BR-3	States of Paraná and São Paulo	BOV	A and H	1	1 August 2008
	BR-4	Part of State of Mato Grosso Do Sul: The area of 15 km from the external borders in the	BOV	A and H	1	1 July 2016]

		municipalit of Porto Murtinho, Caracol, Bela Vista, Antônio João, Ponta Porã, Aral Moreira, Coronel Sapucaia, Paranhos, Sete Quedas, Japorã, and Mundo Novo and the area in the municipalit of Corumbá and Ladário (the former designated highsurveillanc area)	ties				
[F16BW – Botswana	BW-0	Whole country	EQU, EQW				
	BW-1	The veterinary disease control zones 3c, 4b, 5, 8, 9 and 18	BOV, OVI, RUF, RUW	F	1	11 May 2011	26 June 2012
	BW-2	The veterinary disease control zones, 10, 11, 13 and 14	BOV, OVI, RUF, RUW	F	1		7 March 2002

	BW-3	The veterinary disease control zone 12	BOV, OVI, RUF, RUW	F	1	20 October 2008	20 January 2009
	BW-4	The veterinary disease control zone 4a, except the intensive surveillance buffer zone of 10 km along the boundary with the foot-and-mouth disease vaccination zone and wildlife manageme areas	1	F	1	28 May 2013	18 February 2011
	BW-5	The veterinary disease control zone 6, except the intensive surveillanc zone in zone 6 between the border with Zimbabwe and the highway A1	BOV, OVI, RUF, RUW	F	1	28 May 2013	26 June 2012]
BY – Belarus	BY-0	Whole country	_				
BZ – Belize	BZ-0	Whole country	BOV, EQU				
CA – Canada	CA-0	Whole country	BOV, OVI,	G			

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

CH – Switzerland	CH-0 d	Whole	POR, EQU, SUF, SUW, RUF, RUW		
CL – Chile	CL-0	Whole country	BOV, OVI, POR, EQU, RUF, RUW, SUF		
CN – China	CN-0	Whole country	_		
CO – Colombia	CO-0	Whole country	EQU		
CR – Costa Rica	CR-0	Whole country	BOV, EQU		
CU – Cuba	CU-0	Whole country	BOV, EQU		
DZ – Algeria	DZ-0	Whole country	_		
ET – Ethiopia	ET-0	Whole country	_		
FK – Falkland Islands	FK-0	Whole country	BOV, OVI, EQU		
GL – Greenland	GL-0	Whole country	BOV, OVI, EQU, RUF, RUW		
GT – Guatemala	GT-0	Whole country	BOV, EQU		
HK – Hong Kong	HK-0	Whole country	_		
HN – Honduras	HN-0	Whole country	BOV, EQU		

rF3

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^{F3}]						
[^{F4} IL – Israel ^f	IL-0	Whole country	_]
IN – India	IN-0	Whole country	_			
IS – Iceland	IS-0	Whole country	BOV, OVI, EQU, RUF, RUW			
[^{F17} JP — Japan	JP	Whole country	BOV			28 March 2013]
KE – Kenya	KE-0	Whole country	_			
MA – Morocco	MA-0	Whole country	EQU			
ME – Montenegr	ME-0 o	Whole country	BOV, OVI, EQU			
MG – Madagasca	MG-0 r	Whole country	_			
MK – Former Yugoslav Republic of Macedonia	MK-0	Whole country	OVI, EQU			
MU – Mauritius	MU-0	Whole country	_			
MX – Mexico	MX-0	Whole country	BOV, EQU			
NA – Namibia	NA-0	Whole country	EQU, EQW			
	NA-1	South of the cordon fences which extend from Palgrave Point in the west to Gam in the east	BOV, OVI,RUF, RUW	F and J	1	

	Y				1	
NC – New Caledonia	NC-0	Whole country	BOV, RUF, RUW			
NI – Nicaragua	NI-0	Whole country				
NZ – New Zealand	NZ-0	Whole country	BOV, OVI, POR, EQU, RUF, RUW, SUF, SUW			
PA – Panama	PA-0	Whole country	BOV, EQU			
[^{F4} PY – Paraguay	PY-0	Whole country	EQU			
	PY-0	Whole country	BOV	A	1	17 April 2015]
RS – Serbia ^e	RS-0	Whole country	BOV, OVI, EQU			
RU – Russia	RU-0	Whole country				
	RU-1	Region of Murmansk Yamolo- Nenets autonomou area				
[F13SG — Singapore ^g	SG-0	Whole country	NZ- TRANSIT- SG ^h]
SV – El Salvador	SV-0	Whole country	_			
SZ – Swaziland	SZ-0	Whole country	EQU, EQW			
	SZ-1	Area west of the 'red line' fences which extends northwards from the river	BOV, RUF, RUW	F	1	

		Usutu to the frontier with South Africa west of Nkalashane	÷,			
	SZ-2	The veterinary foot and mouth disease surveillanc and vaccination control areas as gazetted as a Statutory Instrument under legal notice number 51 of 2001	1	F	1	4 August 2003
TH – Thailand	TH-0	Whole country	_			
TN – Tunisia	TN-0	Whole country	_			
TR – Turkey	TR-0	Whole country				
	TR-1	The provinces of Amasya, Ankara, Aydin, Balikesir, Bursa, Cankiri, Corum, Denizli, Izmir, Kastamonu Kutahya, Manisa, Usak,	EQU			

		Yozgat and Kirikkale					
UA – Ukraine	UA-0	Whole country	_				
US – United States	US-0	Whole country	BOV, OVI, POR, EQU,SUF, SUW, RUF, RUW	G			
$[^{F18}UY -$	UY-0	Whole	EQU				
Uruguay		country	BOV	A and J	1		1 November 2001
			OVI	A	1]
[F19ZA – South	ZA-0	Whole country	EQU, EQW				
Africa	ZA-1	p o th a a n d c a a s in th v v r o o N n o o o o o o o o o o	BOV, OVI, RUF, RUW he part he cot- nd- nouth lisease ontrol rea ituated he reterinary egions of Journalanga nd Northern provinces, he listrict of ngwavuma	F	1	11 February 2011	

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

		v ro oo N a iii the b a w E e oo le	he border rea vith Botswana ast of ongitude 8°, nd he listrict f Camperdown,
ZW – Zimbabwe	ZW-0	Whole country	

Footnotes:

- a Without prejudice to specific certification requirements provided for in Union agreements with third countries.
- **b** Meat from animals slaughtered on or before the date set out in column 7 may be imported into the Union for 90 days from that date. Consignments carried on vessels on the high seas may be imported into the Union if certified before the date set out in column 7 for 40 days from that date.(N.B.: no date in column 7 means that there are no time restrictions).
- c Only meat from animals slaughtered on or after the date set out in column 8 may be imported into the Union (no date in column 8 means that there are no time restrictions).
- d The former Yugoslav Republic of Macedonia; provisional code that does not prejudge in any way the definitive nomenclature for this country, which will be agreed following the conclusion of negotiations currently taking place on this subject in the United Nations.
- e Not including Kosovo which is at present under international administration pursuant to United Nations Security Council Resolution 1244 of 10 June 1999.
- f [F7Hereafter understood as the State of Israel, excluding the territories under Israeli administration since June 1967, namely the Golan Heights, the Gaza Strip, East Jerusalem and the rest of the West Bank.]
- g [Fi3Only for fresh meat originating from New Zealand, for which New Zealand is authorised for introduction into the Union, which is accompanied by the appropriate model of veterinary certificate issued by the competent authority of New Zealand, destined to the Union and being unloaded, with or without storage and reloaded in an approved establishment during transit through Singapore.
- h Upon entry into the Union, the consignments should be accompanied both by this model of veterinary certificate issued in TRACES by the competent authority of Singapore and by the appropriate model of veterinary certificate for import of fresh meat issued by the competent authority of New Zealand, which may be attached in TRACES by the competent authority of Singapore.]

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

- i [F14For 'RUW': Except from the following departments of the Province of Corrientes: the departments of Berón de Astrada, Capital, Empedrado, General Paz, Itati, Mbucuruyá, San Cosme and San Luís del Palmar.]
- * = Requirements as in accordance with the Agreement between the European Community and the Swiss Confederation on trade in agricultural products (OJ L 114, 30.4.2002, p. 132).
 - = No certificates are laid down and fresh meat imports are prohibited (except for those species where indicated in the line comprising the entry for the whole country).

No offal is authorised for introduction into the Union (except, in the case of bovine species, diaphragm and masseter muscles).]

Textual Amendments

- **F13** Inserted by Commission Implementing Regulation (EU) 2016/535 of 5 April 2016 amending Annex II to Regulation (EU) No 206/2010 as regards the entry of Singapore in the list of third countries, territories or parts thereof from which the introduction into the Union of fresh meat is authorised (Text with EEA relevance).
- F14 Inserted by Commission Implementing Regulation (EU) 2016/922 of 10 June 2016 amending Annex II to Regulation (EU) No 206/2010 as regards the list of third countries, territories or parts thereof from which the introduction into the Union of fresh meat is authorised (Text with EEA relevance).
- F15 Substituted by Commission Implementing Regulation (EU) 2016/922 of 10 June 2016 amending Annex II to Regulation (EU) No 206/2010 as regards the list of third countries, territories or parts thereof from which the introduction into the Union of fresh meat is authorised (Text with EEA relevance).
- **F16** Substituted by Commission Implementing Regulation (EU) No 482/2013 of 24 May 2013 amending Regulation (EU) No 206/2010 laying down lists of third countries, territories or parts thereof authorised for the introduction into the European Union of certain animals and fresh meat and the veterinary certification requirements (Text with EEA relevance).
- **F17** Inserted by Commission Implementing Regulation (EU) No 196/2013 of 7 March 2013 amending Annex II to Regulation (EU) No 206/2010 as regards the new entry for Japan in the list of third countries or parts thereof from which imports into the European Union of certain fresh meat are authorised (Text with EEA relevance).
- F18 Substituted by Commission Implementing Regulation (EU) No 71/2013 of 25 January 2013 amending Regulation (EU) No 206/2010 as regards the entry for Uruguay in the list of third countries, territories or parts thereof authorised for the introduction of fresh meat into the Union and correcting that Regulation as regards the model veterinary certificate for ovine and caprine animals intended for breeding or production after importation (Text with EEA relevance).
- F19 Substituted by Commission Implementing Regulation (EU) No 342/2011 of 8 April 2011 amending Annex II to Regulation (EU) No 206/2010 laying down lists of third countries, territories or parts thereof authorised for the introduction into the European Union of certain animals and fresh meat and the veterinary certification requirements (Text with EEA relevance).

IF20PART 2

Models of veterinary certificates

Model(s):

'BOV'

: Model of veterinary certificate for fresh meat, including minced meat, of domestic bovine animals (including *Bison* and *Bubalus* species and their cross-breeds).

^{&#}x27;1' Category restrictions:

Status.	Point	in tim	a viou	as at	01/07/20	116

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

'OVI' Model of veterinary certificate for fresh meat, including minced meat, of domestic ovine animals (Ovis aries) and domestic caprine animals (Capra hircus). Model of veterinary certificate for fresh meat, including minced meat, 'POR' of domestic porcine animals (Sus scrofa). 'EQU' Model of veterinary certificate for fresh meat, excluding minced meat, of domestic solipeds (Equus caballus, Equus asinus and their cross-'RUF' : Model of veterinary certificate for fresh meat, excluding offal and minced meat, of farmed non-domestic animals of the order Artiodactyla (excluding bovine animals (including *Bison* and *Bubalus* species and their cross-breeds), Ovis aries, Capra hircus, Suidae and Tayassuidae), and of the families Rhinocerotidae and Elephantidae. 'RUW' Model of veterinary certificate for fresh meat, excluding offal and minced meat, of wild non-domestic animals of the order Artiodactyla (excluding bovine animals (including Bison and Bubalus species and their cross-breeds), Ovis aries, Capra hircus, Suidae and Tayassuidae), and of the families Rhinocerotidae and Elephantidae. 'SUF' Model of veterinary certificate for fresh meat, excluding offal and minced meat, of farmed non-domestic animals belonging to the Suidae, Tayassuidae, or Tapiridae families. 'SUW' : Model of veterinary certificate for fresh meat, excluding offal and minced meat, of wild non-domestic animals belonging to the Suidae, Tayassuidae, or Tapiridae families. 'EOW' Model of veterinary certificate for fresh meat, excluding offal and minced meat, of wild solipeds belonging to the subgenus *Hippotigris* (zebra). 'IF13NZ-Model of veterinary certificate only for transit through Singapore with unloading, possible storage and reloading of fresh meat originating from TRANSIT-SG' New Zealand, for which New Zealand is authorised for introduction into the Union, which is eligible for introduction and destined to the Union.] *SG* (Supplementary guarantees) 'A' guarantees regarding the maturation, pH measurement and boning of fresh meat, excluding offal, certified according to the models of veterinary certificates BOV (point II.2.6), OVI (point II.2.6), RUF (point II.2.7) and RUW (point II.2.4). 'C' guarantees regarding the laboratory test for classical-swine-fever in the carcases from which fresh meat was obtained, certified according to the model of veterinary certificate SUW (point II.2.3 B). 'D' guarantees regarding swill feed on holding(s) of animals from which fresh meat certified was obtained according to the model of veterinary certificate POR (point II.2.3 d). 'E' guarantees regarding tuberculosis test in the animals from where fresh meat certified was obtained, according to the model of veterinary certificate BOV (point II.2.4 d). 'F' guarantees regarding the maturation and de-boning of fresh meat, excluding offal, certified according to the models of veterinary certificates BOV (point II.2.6), OVI (point II.2.6), RUF (point II.2.6) and RUW (point II.2.7). 'G' guarantees regarding 1, exclusion of offals and spinal cord; and 2,

testing and origin of cervid animals in relation to chronic wasting

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

disease as referred to in the models of veterinary certificates RUF (point

II.1.7) and RUW (point II.1.8).

: supplementary guarantees required for Brazil. Concerning vaccination

programmes, as the State of Santa Catarina in Brazil does not vaccinate against foot and mouth disease, the reference to a vaccination programme is not applicable for meat coming from animals originating

and slaughtered in that State.

'J' : guarantees regarding the movement of bovine, ovine and caprine

animals from holdings to the slaughterhouse, which allow them to pass via an assembly centre (including markets) before being transported

directly to slaughter.

'[F6K' : holdings or compartments recognised as applying controlled housing

conditions in accordance with Article 8 of Regulation (EC) No

2075/2005.]]

[F20Model BOV]

'H'

Status: Point in time view as at 01/07/2016. **Changes to legislation:** There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

						COUNT	RY
						II.	Health information
						II.1.	Public Health Atte
DUN	TRY			Veterinary certificate to EU			I, the undersigned (EC) No 852/2004, described in Part I
Т	l.1.	Consignor	I.2. Certificate reference No	I.2.a.	_		
		Name	I.O. Combania anno abant anthroite		atio	II.1.1.	the [meat] [minced with Regulation (EC
		Address	I.3. Central competent authority		rtific		(<u>-</u>
₌ │		Tel.	I.4. Local competent authority		Part II: Certification	II.1.2.	the meat has been
dispatched consignment	1.5.	Consignee	1.6.		Part		(1) II.1.3. [the mince internal te
		Name					
De l		Address					II.1.4. the meat has
Date		Postal code					Chapter II of
<u> </u>		Tel.					II.1.5. (1) either [the
rart I: Details of	1.7.	Country of origin ISO code I.8. Region of origin Code	I.9. Country of ISO code destination	I.10. Region of Code destination			A
-	l.11.	Place of origin	l.12.				(¹) <i>or</i> [t
ran		Name Approval number Address					II.1.6. the [meat] [n
		Address					foodstuffs;
Ì	I.13.	Place of loading	I.14. Date of departure				II.1.7. the guarante 96/23/EC, ar
\top	l.15.	Means of transport	I.16. Entry BIP in EU				
		Aeroplane ☐ Ship ☐ Railway wagon ☐					II.1.8. the [meat] [r respectively
		Road vehicle Other	1.17.				
		Identification Documentary references					II.1.9. with regard t
I	I.18.	Description of commodity	I.19. Commodity of	anda (HC anda)			(¹) either [l
			1.19. Commodity of	code (HS code)			(*) eitner [i
			I.	.20. Quantity			
ı	I.21.	Temperature of product	I.	.22. Number of packages			
		Ambient Chilled	Frozen 🗆				
\mid	1.23.	Seal/Container No		.24. Type of packaging			
\perp	1.25.	Commodities certified for:					
		Human consumption					
Ī	1.26.		I.27. For import or admission into	EU 🗆			
+	1.28.	Identification of the commodities	I				
		Species Nature of Treatment (scientific name) commodity type Abai	Approval number of establishments ttoir Cutting plant Cold	packages weight			(¹) or [II.

[F20Model OVI]

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

NTRY	1					Veterinary certific	ate to E
l.1.	Consignor		I.2. Certifica	ite reference No		I.2.a.	
	Name		I.3. Central	competent authori	tv		
	Address						
	Tel.		I.4. Local co	ompetent authority			
1.5.	•		1.6.				
	Name Address						
	Postal code						
	Tel.						
1.7.	Country of origin ISO code	.8. Region of origin Code	I.9. Country destinat		I.10. F	Region of destination	Code
1.11	. Place of origin		I.12.				
	Name Al Address	pproval number					
I.13	. Place of loading		I.14. Date of	departure			
1.15	. Means of transport		I.16. Entry BI	P in EU			
	Aeroplane 🗌 Ship 🔲	Railway wagon					
	Road vehicle Other Identification Documentary references		I.17.				
l.18	. Description of commodity			I.19. Commodity	code (l	HS code)	
					I.20. Q	uantity	
1.21	. Temperature of product				I.22. N	umber of packages	
	Ambient	Chilled	Frozen				
1.23	. Seal/Container No				1.24. Ty	pe of packaging	
1.25	. Commodities certified for:						
	Human consumption						
1.26			I.27. For imp	ort or admission in	to EU		
1.28	. Identification of the commodities						
	Species Nature of	Treatment	Approval numbe	er of establishmen	ts	Number of	Net

COUNTRY П.

Public Health Attest

Health information

I, the undersigned (EC) No 852/2004, caprine animals desc

II.1.1. the [meat] [m accordance w

(1) II.1.2. the meat has

(1) II.1.3. [the minced minternal tempe

II.1.4. the meat has Chapter II of S

II.1.5. (1) either [the Ann

II.1.6. the [meat] [mi foodstuffs;

(1) or

II.1.7. the guarantee 96/23/EC, and

II.1.8. the [meat] [min respectively of

II.1.9. with regard to

(1) either [II.1.9.1. for imp

(b) t

(1) [(c) if

[II.1.9.2. for in (1) or

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

Model POR

	CO	UNTRY	Veterinary certificate to EU			
	l.1.	Consignor	I.2. Certificate reference number I.2.a.			
		Name	I.3. Central Competent Authority			
		Address				
ent		Tel. No	I.4. Local Competent Authority			
gnm	1.5.	Consignee	1.6.			
onsi		Name				
o pe		Address				
tche		Postal code				
ispa		Tel. No				
Part I: Details of dispatched consignment	1.7.	Country ISO I.8. Region Code of origin of origin	I.9. Country of ISO I.10. Region of Code destination code destination			
Det	1.11.	. Place of origin	1.12.			
ırı		Name Approval number				
ß.		Address				
	I.13	. Place of loading	I.14. Date of departure			
	1.15	. Means of transport	I.16. Entry BIP in EU			
		Aeroplane Ship Railway wagon				
		Road vehicle Other				
		Identification:	1.17.			
		Documentary references:				
	I.18	. Description of commodity	I.19. Commodity code (HS code)			
			I.20. Quantity			
	1.21	. Temperature of product	I.22. Number of packages			
		Ambient Chiled	Frozen			
	1.23	l. Identification of container/seal number	I.24. Type of packaging			
	1.25	. Commodities certified for:	'			
		Human consumption				
	1.26		I.27. For import or admission into EU			
	1.28	s. Identification of the commodities				
	(\$	Scientific name) commodity type	oroval number establishments Number Net of packages weight			
		Abatto	ir Cutting plant Cold store			

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model POR

	П.	Health	h information		II.a. Certificate reference number	II.b.			
	II.1.	Public	Health Attesta	tion					
I, the undersigned official veterinarian, declare that I am aware of the relevant requirements of Regulations (EC) (EC) No 852/2004, (EC) No 853/2004 and (EC) No 854/2004 and hereby certify that the meat of domestic sw in Part I was produced in accordance with those requirements, in particular that:									
fication	II.1.1 the [meat] [minced meat] (¹) comes from (an) establishment(s) implementing a programme based of principles in accordance with Regulation (EC) No 852/2004;								
Part II: Certification		II.1.2	the meat has b No 853/2004;	een obtai	ned in compliance with the conditions set out	in Section I of Annex III to Regulation (EC)			
Par	▶ ⁽¹⁾	II.1.3	II.1.3 the meat fulfils the requirements of Regulation (EC) No 2075/2005 laying down specific rules on official control <i>Trichinella</i> in meat, and in particular:						
			(1) either	[has bee	n subjected to an examination by a digestion r	nethod with negative results;]			
			(¹) or	[has bee 2075/20	en subjected to a freezing treatment in accord5;]	dance with Annex II to Regulation (EC) No			
	(¹)(²) or [is derived from domestic porcine animals either coming from a holding officially recognise plying controlled housing conditions in accordance with Article 8 of Regulation (EC) No 20 or not weaned and less than 5 weeks of age.] ◀								
	(¹) II.1.4 [the minced meat has been produced in accordance with Section V of Annex III to Regulation (EC) No 853/2004 a frozen to an internal temperature of not more than –18 °C;]								
		II.1.5 the meat has been found fit for human consumption following ante and post-mortem inspections carri accordance with Chapter II of Section I and Chapters IV and IX of Section IV of Annex I to Regulat No 854/2004;							
					arcass or parts of the carcass have been marked with a health mark in accordance with ter III of Section I of Annex I to Regulation (EC) No 854/2004;]				
			(¹) or		packages of [meat] [minced meat] (1) have been marked with an identification mark in dance with Section I of Annex II to Regulation (EC) No 853/2004;]				
		II.1.7	the [meat] [mine criteria for food		(¹) satisfies the relevant criteria set out in Regul	lation (EC) No 2073/2005 on microbiological			
		II.1.8			live animals and products thereof provided by and in particular Article 29, are fulfilled.	the residue plans submitted in accordance			
		II.1.9			t] (¹) has been stored and transported in acc vely of Annex III to Regulation (EC) No 853/20				
	(²) [I	(²) [II.1.10 it fulfils the requirements of Regulation (EC) No 1688/2005 implementing Regulation (EC) No 853/2004 as regar special guarantees concerning Salmonella for consignments to Finland and Sweden of certain meat and eggs;]							
	II.2.	Anima	l Health attesta	tion					
		I, the u	ndersigned offic	ial veterina	arian, hereby certify, that the fresh meat descri	bed in Part I :			
		II.2.1	has been obtai	ned in the	territory/ies with code:(3)	which, at the date of issuing this certificate:			
			(¹) either		been free for 12 months from foot-and-mout sical swine fever, swine vesicular disease, and]				
			(¹) or		nas been free for 12 months from rinderpest, Afric classical swine fever] (') and [swine vesicular d				

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model POR

II.	Heal	Health information			II.a. Certificate reference number	II.b.
					has been considered free from [foot-and-mout [swine vesicular disease] (¹), sincehad cases/outbreaks afterwards, and author Regulation (EC) No/, of	(dd/mm/yyyy), without having rised to export this meat by Commission(dd/mm/yyyy), and]
				imp	ng the last 12 months no vaccination against orts of domestic animals vaccinated against tory;	
		11.2.2	has been obtai	ned from	animals that:	
			(¹) either		mained in the territory described under point l before slaughter;]	I.2.1 since birth, or for at least the last three
			(¹) or	point II.2	een introduced on(dd/ 2.1, from the territory with code	
			(¹) or		een introduced on(dd/ 2.1, from the EU Member State	
		II.2.3	has been obtai	ned from	animals coming from holdings:	
			(a) in which r point II.2.1		ne animals present therein have been vacci	inated against the diseases referred to in
					, in an area of 10 km radius, there has been no e previous 40 days,	case/outbreak of the diseases referred to in
			(c) that are no weeks;	t subject	to prohibition as a result of an outbreak of	porcine brucellosis during the previous six
		(1) (4)			g has been received that pigs are not fed with on the list established by the competent authority for the competen	
		11.2.4	has been obtai	ned from	animals that:	
			(a) have remain	ned sepa	rate since birth from wild cloven-hoofed anima	ls,
				ouse with	ed from their holdings in vehicles, cleaned and out contact with other animals which did not com	
					e, have passed ante-mortem health inspection wn no evidence of the diseases referred to in po	
					red on(dd/mm/yyyy) or b (dd/mm/yyyy). (5);	petween (dd/mm/yyyy)
		II.2.5	of the disease preparation of	s referred meat for i	n establishment around which, within a radius I to in point II.2.1 during the previous 40 days mportation into the Union has been authorised If the total cleaning and disinfection of the es	s or, in the event of a case of disease, the d only after slaughter of all animals present,
		II.2.6	has been obtai certificate.	ned and p	orepared without contact with other meats not c	complying with the conditions required in this
▶ ⁽¹⁾	II.3.	Anima	I welfare attest	ation		
		mals w evant p	hich have been l	nandled ir n legislat	arian, hereby certify, that the fresh meat describe the slaughterhouse before and at the time of s ion and have met requirements at least equivale /2009 (⁶). ◀	slaughter or killing in accordance with the rel-

Status: Point in time view as at 01/07/2016.

COU	ITR	Y		Model PC					
II.		Health information	II.a. Certificate reference number	II.b.					
	No	tes							
	Thi	s certificate is meant for fresh meat, inclu	uding minced meat, of domestic swine (Sus	scrofa).					
	Fresh meat means all animal parts fit for human consumption whether fresh, chilled or frozen.								
	Part I:								
	_	Box reference I.8: Provide the code of te	erritory as appearing in Part 1 of Annex II to	Regulation (EU) No 206/2010.					
	_	Box reference I.11: Place of origin: name	e and address of the dispatch establishmen	t.					
	_		r (railway wagons or container and lorries), ading, the consignor must inform the BIP of o	flight number (aircraft) or name (ship) is to be entry into the Union.					
	_	Box reference I.19: Use the appropriate	HS code: 02.03, 02.06, 02.09, 05.04 or 15.0	01,					
	_	Box reference I.20: Indicate total gross	weight and total net weight.						
	_	Box reference I.23: For containers or bo	xes, the container number and the seal num	ber (if applicable) should be included.					
	_	Box reference I.28: Nature of commodity	y: Indicate 'carcass-whole', 'carcass-side', 'c	arcass-quarters', 'cuts' or 'minced meat'.					
		Minced meat is deboned meat that has muscle (including the adjoining fatty tiss		have been prepared exclusively from striated					
	-	Box reference I.28: Treatment type: If apport freezing (mm/yy) of the cuts/pieces.	propriate, indicate 'deboned'; 'bone in'; 'matu	red' and/or 'minced'. If frozen, indicate the date					
	Pai	rt II:							
	(¹)	Keep as appropriate.							
	(2)	Delete if the consignment is not intende	d for import into Finland or Sweden.						
	(3)	Code of the territory as it appears in Par	rt 1 of Annex II to Regulation (EU) No 206/20	010.					
	(4)	Supplementary guarantees to be provide with the entry 'D'.	ded when required in column 5 'SG' of Part	1 of Annex II to Regulation (EU) No 206/2010,					
		Catering waste means: all waste from foo industrial kitchens and household kitchen		aurants, catering facilities or kitchens, including					
		of authorisation for importation into the liperiod where restrictive measures have part thereof.	Union of the third country, territory or part the	om animals slaughtered either prior to the date ereof referred to in boxes I.7 and I.8, or during a of this meat from this third country, territory or					
		OJ L 303, 18.11.2009, p. 1. ◀							
▶ ⁽²⁾	(7)	Only for third countries with the entry 'K	' in column 'SG' in Part 1 of Annex II to Reg	ulation (EU) No 206/2010. ◀					
	Off	icial veterinarian							
		Name (in capital letters):	Qualificati	on and title:					
		Date:	Signature	:					
		Stamp:							

Status: Point in time view as at 01/07/2016. **Changes to legislation:** There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

Model EQU

	CO	UNTRY			veterinary certificate to EU		
	1.1.	Consignor	I.2. Certificate r	eference nun	nber I.2.a.		
		Name	I.3. Central Con	npetent Autho	prity		
		Address	I.4. Local Comp		-		
ent		Tel. No	1.4. Local Comp	eterit Autrior	nty		
gum	I.5.	Consignee	1.6.				
onsi		Name					
pe		Address					
tch		Postal code					
lispa		Tel. No			T		
Part I: Details of dispatched consignment	1.7.	Country ISO I.8. Region Code of origin of origin	I.9. Country of destination	ISO code	I.10. Region of Code destination		
Det	1.11.	. Place of origin	l.12.				
art :		Name Approval number Address					
ă		Address					
	1.13	. 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					
		Identification:	l.17.				
		Documentary references:					
	I.18	. Description of commodity	1.19	9. Commodit	y code (HS code)		
				1.:	20. Quantity		
	1.21	. Temperature of product		Li	22. Number of packages		
		Ambient Chiled	Frozen				
	1.23	l. Identification of container/seal number		1.3	24. Type of packaging		
	1.25	Commodities certified for:					
		numan consumption					
	1.26		I.27. For import o	r admission i	nto EU		
	1.28	s. Identification of the commodities					
	(\$	Species Nature of Approval n Scientific name) commodity	umber establishme	nts	Number Net of packages weight		
		Abattoir C	Cutting plant Col	d store			

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model EQU

	II.	Health	information		II.a. Certificate reference number	II.b.				
	II.1.	Public	Health Attestat	tion						
		(EC) N	o 852/2004, (EC) No 853/2		an, declare that I am aware of the relevant requirements of Regulations (EC) No 178/2002, 04 and (EC) No 854/2004 and hereby certify that the meat of domestic solipeds described be with those requirements, in particular that:				
Part II: Certification		II.1.1 the meat comes from (an) establishment(s) implementing a programme based on the HACCP accordance with Regulation (EC) No 852/2004;								
t II: Cerl		II.1.2	the meat has b No 853/2004;	een obtai	ned in compliance with the conditions set out	in Section I of Annex III to Regulation (EC)				
Par	II.1.3 the meat fulfils the requirements of Regulation (EC) No 2075/2005 laying down specific rules for Trichinella in meat, and in particular, has been subject to an examination by a digestion me results;									
		II.1.4			d fit for human consumption following ante a er II of Section I and Chapters III and IX of					
		II.1.5	(¹) either		ass or parts of the carcass have been mark III of Section I of Annex I to Regulation (EC) No					
(¹) or [the packages of meat have been marked with an identification mark in accordan Annex II to Regulation (EC) No 853/2004;]										
		II.1.6	the meat satisfoodstuffs;	fies the re	elevant criteria set out in Regulation (EC) No	o 2073/2005 on microbiological criteria for				
		II.1.7			live animals and products thereof provided by and in particular Article 29 thereof, are fulfilled;					
		II.1.8	the meat has been stored and transported in accordance with the relevant requirements of Section I of An Regulation (EC) No 853/2004.							
	II.2.	Anima	l Health attesta	tion						
		I, the u	ndersigned offici	ial veterina	arian, hereby certify, that the fresh meat descri	bed in Part I:				
		II.2.1	has been obtai	ned in the	territory/ies with code:	(2);				
		11.2.2	has been obtain	ned from o	domestic solipeds, which:					
			(¹) either		nained in the territory described under point II before slaughter;]	.2.1 since birth, or for at least the last three				
		(¹) or [have been introduced on								
			(¹) or		en introduced on(dd/i .1, from the EU Member State					
		II.2.3	which, within a previous 40 day has been autho	radius of ys or, in th orised onl	animals which were slaughtered on	d/mm/yyyy) (³) in a slaughterhouse around rican horse sickness or glanders during the ration of meat for importation into the Union oval of all meat, and the total cleaning and				

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY	Model EQU
CONTIN	Model Edo

П. Health information II.a. Certificate reference number II.b.

> has been obtained and prepared without contact with other meats not complying with the conditions required in this certificate

▶⁽¹⁾ Ⅱ.3. Animal welfare attestation

I, the undersigned official veterinarian, hereby certify, that the fresh meat described in Part I of this certificate derives from animals which have been handled in the slaughterhouse before and at the time of slaughter or killing in accordance with the relevant provisions of Union legislation and have met requirements at least equivalent to those laid down in Chapters II and III of Council Regulation (EC) No 1099/2009 (4). ◀

Notes

This certificate is meant for fresh meat, excluding minced meat, of domestic solipeds (Equus caballus, Equus asinus and their crossbreeds).

Fresh meat means all animal parts fit for human consumption whether fresh, chilled or frozen.

Part I:

- Box reference I.8: Provide the code of territory as appearing in Part 1 of Annex II to Regulation (EU) No 206/2010.
- Box reference I.11: Place of origin: name and address of the dispatch establishment.
- Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor must inform the BIP of entry into the Union.
- Box reference I.19: Use the appropriate HS code: 02.05, 02.06 or 05.04.
- Box reference I.20: Indicate total gross weight and total net weight.
- Box reference I.23: For containers or boxes, the container number and the seal number (if applicable) should be included.
- Box reference I.28: Nature of commodity: Indicate 'carcass-whole', 'carcass-side', 'carcass-quarters' or 'cuts'.
- Box reference I.28: Treatment type: If appropriate, indicate 'deboned'; 'bone in' and/or 'matured'. If frozen, indicate the date of freezing (mm/yy) of the cuts/pieces.

Part II:

- (1) Keep as appropriate.
- (2) Code of the territory as it appears in Part 1 of Annex II to Regulation (EU) No 206/2010.
- (3) Dates: imports of this meat shall not be authorised when obtained from animals slaughtered either prior to the date of authorisation

	()	for importation into the Union of the third country, territory or part thereof referred to in boxes I.7 and I.8, or during a period where restrictive measures have been adopted by the Union against imports of this meat from this third country, territory or part thereof.						
▶ ⁽²⁾	(4)	OJ L 303, 18.11.2009, p. 1. ◀						
	Off	icial veterinarian						
		Name (in capital letters):	Qualification and title:					
		Date:	Signature:					
		Stamp:						

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

Model RUF

	CO	UNTRY	Veterinary certificate to EU		
	1.1.	Consignor	I.2. Certificate reference number I.2.a.		
		Name	I.3. Central Competent Authority		
		Address			
ent		Tel. No	I.4. Local Competent Authority		
n n	1.5.	Consignee	1.6.		
nsić		Name			
oo p		Address			
tche		Postal code			
spa		Tel. No			
Part I: Details of dispatched consignment	1.7.	Country ISO I.8. Region Code of origin code	I.9. Country of ISO I.10. Region of Code destination code		
Deta	l.11.	. Place of origin	1.12.		
ī.		Name Approval number			
Ра		Address			
	I.13.	. Place of loading	I.14. Date of departure		
	1.15.	. Means of transport	I.16. Entry BIP in EU		
		Aeroplane Ship Railway wagon			
		Road vehicle Other			
		Identification:	I.17.		
		Documentary references:			
	I.18	. Description of commodity	I.19. Commodity code (HS code)		
			I.20. Quantity		
	1.21	. Temperature of product	I.22. Number of packages		
		Ambient Chiled	Frozen		
	1.23	. Identification of container/seal number	I.24. Type of packaging		
	1.25	Commodities certified for: Human consumption			
	1.26		I.27. For import or admission into EU		
	1.28	d. Identification of the commodities			
	(\$	Species Nature of Treatment App Scientific name) commodity type Abattoi	roval number establishments Number Net of packages weight r Cutting plant Cold store		

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model RUF

II. Health information II.a. Certificate reference number II.b. **Public Health Attestation** II.1. I, the undersigned official veterinarian, declare that I am aware of the relevant requirements of Regulations (EC) No 178/2002, (EC) No 852/2004, (EC) No 853/2004, (EC) No 854/2004 and (EC) No 999/2001 and hereby certify that the meat of farmed animals of the order Artiodactyla (excluding bovine animals (including Bison and Bubalus species and their cross-breeds), Ovis aries, Capra hircus, Suidae and Tayassuidae), and of the families Rhinocerotidae and Part II: Certification Elephantidae described in Part I was produced in accordance with those requirements, in particular that: II.1.1 the meat comes from (an) establishment(s) implementing a programme based on the HACCP principles in accordance with Regulation (EC) No 852/2004; II.1.2 the meat has been obtained in accordance with the conditions set out in Section III of Annex III to Regulation (EC) No 853/2004; II.1.3 the meat has been found fit for human consumption following ante and post-mortem inspections carried out in accordance with Chapter II of Section I and Chapters VII and IX of Section IV of Annex I to Regulation (EC) No 854/2004; II.1.4 (1) either [the carcass or parts of the carcass have been marked with a health mark in accordance with Chapter III of Section I of Annex I to Regulation (EC) No 854/2004;] (1) or [the packages of meat have been marked with an identification mark in accordance with Section I of Annex II to Regulation (EC) No 853/2004;] II.1.5 the meat satisfies the relevant criteria set out in Regulation (EC) No 2073/2005 on microbiological criteria for foodstuffs: the guarantees covering live animals and products thereof provided by the residue plans submitted in accordance II.1.6 with directive 96/23/EC, and in particular Article 29 thereof, are fulfilled (1) (2) [II.1.7 with regard to Chronic Wasting Disease (CWD): This product contains or is derived exclusively from meat, excluding offal and spinal cord, of farmed cervid animals which have been examined for Chronic Wasting Disease by histopathology, immunohistochemistry or other diagnostic method recognised by the competent authority with negative results and is not derived from animals coming from a herd where Chronic Wasting Disease has been confirmed or is officially suspected.] the meat has been stored and transported in accordance with the relevant requirements of Section I of Annex III to Regulation (EC) No 853/2004. 11.2. **Animal Health attestation** I, the undersigned official veterinarian, hereby certify, that the fresh meat described in Part I: (a) has been free for 12 months from rinderpest, and during the same period no vaccination against this disease has taken place, and (1) either [(b) has been free for 12 months from foot-and-mouth disease, and during the same period no vaccination against this disease has taken place;] [(b) has been considered free from foot-and-mouth disease since .. (1) or .. (dd/mm/yyyy), without having had cases/outbreaks afterwards, and authorised to export this meat by Commission Regulation (EU) No [(b) vaccination programmes against foot-and-mouth disease are being officially carried out and controlled in (1) (4) or domestic bovine animals;)

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model RUF

II.	Health	information	II.a. Certificate reference number	II.b.
	II.2.2 has been obtained from a		animals that:	
			mained in the territory described under point II before slaughter;]	.2.1 since birth, or for at least the last three
		point II.2	een introduced on(dd/r 2.1, from the territory with code t this fresh meat into the Union;]	
	II.2.3	has been obtained from	animals coming from holdings:	
		(a) in which none of to or] (5) rinderpest,	the animals present therein have been vac	ccinated against [foot-and-mouth disease
			nary inspections are carried out to diagnose dia are not subject to prohibition as a result of an ou	
	(¹) either	[(c) in and around which rinderpest during the	in an area of 10 km radius, there has been no or previous 30 days,]	case/outbreak of foot-and-mouth disease or
	(¹) (⁴) or		icial restriction for health reasons and in and arc utbreak of foot-and-mouth disease or rinderpes	
		(d) where the animals ha	ave remained for at least 40 days before direct	dispatch to the slaughterhouse;]
	II.2.4	has been obtained from	animals:	
	(¹) either		ansported from their holdings in vehicles, clea ouse, without contact with other animals which o	
			erhouse, have passed ante-mortem health insp ve shown no evidence of the diseases referred	
			ughtered on(dd/mm/ (dd/mm/yyyy) (⁶);]	/yyyy) or between
	(¹) or		laughtered on the holding of origin, following olding, who has provided a written statement	
			unacceptable risk would have been posed to thof the animals to an slaughterhouse,	ne welfare of the animals or to their handlers
	animals, — the animals have in particular, hav — the animals wer (dd/mm/yyyy), (been inspected and authorised by the comp	petent authority for the slaughter of game
			e passed the ante-mortem health inspection du e shown no evidence of the diseases referred t	
			e slaughtered between)	. (dd/mm/yyyy) and
			he animals was performed correctly, and	
		 the slaughtered 	animals were eviscerated within three hours of	the time of slaughter, and
		where more than one	ch have been transported to the approved slau e hour elapsed since the time of slaughter, a te rrival of the vehicle used for the transport;]	
	(¹) (²) II.2.5	[has been obtained from	animals that have remained since birth or for the	he last 3 months separate from wild cloven-

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model RUF

II. Health informati	ion	II.a. Certificate reference number	II.b.
II.2.6	of the diseases r preparation of me	d in an establishment around which, within a rad ferred to in point II.2.1 during the previous 30 of at for importation into the Union has been author at, and the total cleaning and disinfection of the	days or, in the event of a case of disease, the rised only after slaughter of all animals present,
II.2.7			
		as been obtained and prepared without contact wi quired above.]	ith other meats not complying with the conditions
	C Si	ontains boneless meat, obtained only from de-bo reasses in which the main accessible lymphatic bmitted to maturation at a temperature above + 2 moved and in which the pH value of the meat v iddle of the longissimus-dorsi muscle after matur	c glands have been removed, which have been 2 °C for at least 24 hours before the bones were was below 6.0 when tested electronically in the
	С	is been kept strictly separate from meat not c intificate during all stages of its production, de- tixes or cartons for further storage in dedicated ar	coning and storage until it has been packed in
	C S	ontains boneless meat, obtained only from de-bo rcasses in which the main accessible lymphatic bmitted to maturation at a temperature above + 2 moved, and	glands have been removed, which have been
		is been kept strictly separate from meat not c	conforming to the requirements set out in this poning and storage until it has been packed in

boxes or cartons for further storage in dedicated areas.]

▶⁽¹⁾ (¹) II.3. Animal welfare attestation

In case the fresh meat described in Part I of this certificate derives from animals which have been slaughtered or killed in a slaughterhouse, I, the undersigned official veterinarian, hereby certify, that they were handled in the slaughterhouse before and at the time of slaughter or killing in accordance with the relevant provisions of Union legislation and have met requirements at least equivalent to those laid down in Chapters II and III of Council Regulation (EC) No 1099/2009 (°). ◀

Notes

This certificate is meant for fresh meat, excluding offal and minced meat, of wild animals of the order Artiodactyla (excluding bovine animals (including *Bison* and *Bubalus* species and their cross-breeds), *Ovis aries, Capra hircus*, Suidae and Tayassuidae), and of the families Rhinocerotidae and Elephantidae, that are domestically kept or bred since birth or for the last three months in farms.

Fresh meat means all animal parts fit for human consumption whether fresh, chilled or frozen.

Part I:

- Box reference I.8: Provide the code of territory as appearing in Part 1 of Annex II to Regulation (EU) No 206/2010.
- Box reference I.11: Place of origin: name and address of the dispatch establishment.
- Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor must inform the BIP of entry into the Union.
- Box reference I.19: Use the appropriate HS code: 02.06, 02.08.90 or 05.04.
- Box reference I.20: Indicate total gross weight and total net weight.
- Box reference I.23: For containers or boxes, the container number and the seal number (if applicable) should be included.
- Box reference I.28: Nature of commodity: Indicate 'carcass-whole', 'carcass-side', 'carcass-quarters', or 'cuts'.
- Box reference I.28: Treatment type: If appropriate, indicate 'deboned'; 'bone in' and/or 'matured'. If frozen, indicate the date of freezing (mm/yy) of the cuts/pieces.

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model RUF

II.		Health information	II.a. Certificate reference number	II.b.					
	Part II:								
		Keep as appropriate. Supplementary guarantees regarding fresh meat obtained from cervids to be provided when required in column 5 'SG' of Part 1 of Annex II to Regulation (EU) No 206/2010, with the entry 'G'.							
	(3)	Code of the territory as it appears in Par	10.						
	(4)	Supplementary guarantees regarding Part 1 of Annex II to Regulation (EU) N	provided when required in column 5 'SG' of						
(5) Delete when the exporting country carries out vaccination against foot-and-mouth disease with serotypes A, O or C country is allowed for import into the Union matured de-boned meat which fulfils the supplementary guarantees descrit footnote (4).									
	(⁶)	date of authorisation for importation into	the Union of the third country, territory or p	d from animals slaughtered either prior to the art thereof referred to in boxes I.7 and I.8, or it imports of this meat from this third country,					
		Not necessary for farmed game animals							
	(⁸)		010, with the entry 'F'. The matured de-bone	ided when required in column 5 'SG' of Part 1 d meat shall not be authorised for importation					
▶ ⁽¹⁾	(9)	OJ L 303, 18.11.2009, p. 1. ◀							
	Off	icial veterinarian							
		Name (in capital letters):	Qualification	n and title:					
		Date:	Signature:						
		Stamp:							

Status: Point in time view as at 01/07/2016. **Changes to legislation:** There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

Model RUW

	CO	COUNTRY Veterinary certificate to EU					
	1.1.	Consignor	I.2. Certificate reference number I.2.a.				
		Name	I.3. Central Competent Authority				
		Address	, ,				
ent		Tel. No	I.4. Local Competent Authority				
gnm	1.5.	Consignee	1.6.				
onsi		Name					
o pe		Address					
tche		Postal code					
ispa		Tel. No					
Part I: Details of dispatched consignment	1.7.	Country ISO I.8. Region Code of origin code	I.9. Country of ISO I.10. Region of Code destination code destination				
Det	1.11.	. Place of origin	1.12.				
ırt I:		Name Approval number					
Pe		Address					
	1.13	. 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					
		Identification:	1.17.				
		Documentary references:					
	I.18	. Description of commodity	I.19. Commodity code (HS code)				
			I.20. Quantity				
	1.21	. Temperature of product	I.22. Number of packages				
		Ambient Chiled Chiled	Frozen				
	1.23	l. Identification of container/seal number	I.24. Type of packaging				
	1.25	. Commodities certified for:	·				
		Human consumption					
	1.26		I.27. For import or admission into EU				
	1.28	s. Identification of the commodities					
	(\$	Species Nature of Treatment App Scientific name) commodity type	roval number establishments Number Net of packages weight				
		Abatto	ir Cutting plant Cold store				
I							

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model RUW

II. Health information II.a. Certificate reference number II.b. **Public Health Attestation** II.1. I, the undersigned official veterinarian, declare that I am aware of the relevant requirements of Regulations (EC) No 178/2002, (EC) No 852/2004, (EC) No 853/2004 and (EC) No 854/2004 and hereby certify that the fresh meat of wild animals of the order Artiodactyla (excluding bovine animals (including Bison and Bubalus species and their cross-breeds), Ovis aries, Capra hircus, Suidae and Tayassuidae), and of the families Rhinocerotidae and Elephantidae described in Part II: Certification Part I was produced in accordance with those requirements, in particular that: 11.1.1 the meat comes from (an) establishment(s) implementing a programme based on the HACCP principles in accordance with Regulation (EC) No 852/2004; II.1.2 the meat has been obtained in compliance with the conditions set out in Section IV of Annex III to Regulation 853/2004, and in particular: (i) before skinning, it has been stored and handled separately from other food and not frozen; and (ii) after skinning, it has undergone a final inspection as referred to in point II.1.4; (1) II.1.3 [in the case of susceptible species, the meat fulfils the requirements of Regulation (EC) No 2075/2005 laying down specific rules on official controls for Trichinella in meat;] the meat has been found fit for human consumption following a post-mortem inspection carried out in accordance with Chapter II of Section I and Chapters VIII and IX of Section IV of Annex I to Regulation (EC) No 854/2004; II.1.5 (1) either [in the case of large wild game, the carcass or parts of the carcass have been marked with a health mark in accordance with Chapter III of Section I of Annex I to Regulation (EC) No 854/2004;] [the packages of meat have been marked with an identification mark in accordance with Section I of (1) or Annex II to Regulation (EC) No 853/2004;] the meat satisfies the relevant criteria set out in Regulation (EC) No 2073/2005 on microbiological criteria for foodstuffs; 11.1.7 the guarantees covering live animals and products thereof provided by the residue plans submitted in accordance with Directive 96/23/EC, and in particular Article 29 thereof, are fulfilled (1) (2) [II.1.8 with regard to Chronic Wasting Disease (CWD): This product contains or is derived exclusively from meat, excluding offal and spinal cord, of wild cervid animals which have been examined for Chronic Wasting Disease by histopathology, immunohistochemistry or other diagnostic method recognised by the competent authority with negative results and is not derived from animals coming from a region where Chronic Wasting Disease has been confirmed in the last three years or is officially suspected. the meat has been stored and transported in accordance with the relevant requirements of Section I of Annex III to Regulation (EC) No 853/2004. 11.2. Animal Health attestation I, the undersigned official veterinarian, hereby certify, that the fresh meat described in Part I: (a) has been free for 12 months from rinderpest, and during the same period no vaccination against this disease has taken place, and (1) either (b) has been free for 12 months from foot-and-mouth disease, and during the same period no vaccination against

this disease has taken place;]

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model RUW

II. Health information		II.a. Certificate reference number	II.b.
having had cases/ou		considered free from foot-and-mouth disease since cases/outbreaks afterwards, and authorised to exp/, of	
(¹) (⁴) or		programmes against foot-and-mouth disease are ovine animals;]	being officially carried out and controlled in
II.2.2		ned from wild animals that were killed between (dd/mm/yyyy) (5) inside the territory referre	
	` '	e that exceeds 20 km from the borders of a country or porting this fresh meat into the Union,	part thereof, which is not authorised during this
	(b) in an area point II.2.1;	where during the last 60 days, there has been r	o restrictions for the diseases referred to in
II.2.3	game-handling diseases referre of meat for impo	ed from animals which after killing were transported establishment around which, within a radius of 10 d to in point II.2.1 during the previous 30 days or, in tration into the Union has been authorised only after e establishment under the control of an official veter	km, there has been no case/outbreak of the the event of a case of disease, the preparation removal of all meat, and the total cleaning and
II.2.4			
	(¹) either	[has been obtained and prepared without contact wit required above.]	n other meats not complying with the conditions
carcasse submitte removed		[contains boneless meat, obtained only from de-bon carcasses in which the main accessible lymphatic submitted to maturation at a temperature above +2 removed and in which the pH value of the meat w middle of the longissimus-dorsi muscle after matura	glands have been removed, which have been °C for at least 24 hours before the bones were as below 6.0 when tested electronically in the
		has been kept strictly separate from meat not co- certificate during all stages of its production, de-b- boxes or cartons for further storage in dedicated are	oning and storage until it has been packed in
	(¹) (⁶) or	[contains boneless meat, obtained only from de-bon- carcasses in which the main accessible lymphatic submitted to maturation at a temperature above +2 removed, and	glands have been removed, which have been
certificat		has been kept strictly separate from meat not co certificate during all stages of its production, de-b- boxes or cartons for further storage in dedicated are	oning and storage until it has been packed in

Notes

This certificate is meant for fresh meat, excluding offal and minced meat, of wild animals of the order Artiodactyla (excluding bovine animals (including *Bison* and *Bubalus* species and their cross-breeds), *Ovis aries, Capra hircus*, Suidae and Tayassuidae), and of the families Rhinocerotidae and Elephantidae that are killed or hunted in the wild.

Fresh meat means all animal parts fit for human consumption whether fresh, chilled or frozen.

After importation, unskinned carcasses must be conveyed without delay to the processing establishment of destination.

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model RUW

II.	Health information	II.a. Certificate reference number	II.b.				
Pa	Part I:						
_	Box reference I.8: Provide the code of to	erritory as appearing in Part 1 of Annex II to Re	egulation (EU) No 206/2010.				
_		e and address of the dispatch establishment.	•				
_		er (railway wagons or container and lorries), fli ading, the consignor must inform the BIP of en					
—	Box reference I.19: Use the appropriate	HS code: 02.01, 02.02, 02.04, 02.06, 02.08.9	0 or 05.04.				
—	Box reference I.20: Indicate total gross	weight and total net weight.					
—	Box reference I.23: For containers or bo	oxes, the container number and the seal numb	er (if applicable) should be included.				
—	Box reference I.28: Nature of commodit	ty: Indicate 'carcass-whole', 'carcass-side', 'car	cass-quarters' or 'cuts'.				
_	Box reference I.28: <i>Treatment type</i> : If an of the cuts/pieces.	opropriate, indicate 'matured' or 'unskinned'. If	frozen, indicate the date of freezing (mm/yy)				
-	Box reference I.28: Abattoir: any abatto	ir or game handling establishment.					
Pa	rt II:						
(¹)	Keep as appropriate						
(2)	Supplementary guarantees regarding of Annex II to Regulation (EU) No 206	fresh meat obtained from cervids to be provid 5/2010, with the entry ' G '.	ed when required in column 5 'SG' of Part 1				
(3)	Code of the territory as it appears in Pa	rt 1 of Annex II to Regulation (EU) No 206/201	0.				
(4)	Supplementary guarantees regarding Part 1 of Annex II to Regulation (EU) I	meat from matured de-boned meat to be p No 206/2010 with the entry 'A'.	rovided when required in column 5 'SG' of				
	The matured de-boned meat shall not animals.	be authorised for importation into the Union	until 21 days after the date of killing of the				
(5)	for importation into the Union of the thir	uthorised when obtained from animals killed or rd country, territory or part thereof referred to i d by the Union against imports of this meat fro	n boxes I.7 and I.8, or during a period where				
(⁶)	Supplementary guarantees regarding meats from matured de-boned meat to be provided when required in column 5 'SG' of Part 1 of Annex II to Regulation (EU) No 206/2010, with the entry 'F'. The matured de-boned meat shall not be allowed for importation into the Union until 21 days after the date of slaughter of the animals.						
	the official and 21 days after the date of	slaughter of the arimals.					
Off	Official veterinarian						
	Name (in capital letters): Qualification and title:						
	Date: Signature:						
	Stamp:						
	-						

Status: Point in time view as at 01/07/2016. **Changes to legislation:** There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

Model SUF

	COUNTRY Veterinary certificate to EU						
	1.1.	Consignor	I.2. Certificate reference number I.2.a.				
		Name	I.3. Central Competent Authority				
		Address	I.4. Local Competent Authority				
ent		Tel. No	1.4. Local competent Admonty				
gnm	1.5.	Consignee	1.6.				
onsi		Name					
o pa		Address					
atch		Postal code					
dsip		Tel. No					
Part I: Details of dispatched consignment	1.7.	Country ISO I.8. Region Code of origin of origin	I.9. Country of ISO I.10. Region of Code destination code destination				
Det	1.11.	. Place of origin	1.12.				
art		Name Approval number Address					
۵		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					
		Identification:	1.17.				
		Documentary references:					
	I.18	. Description of commodity	I.19. Commodity code (HS code)				
			I.20. Quantity				
	1.21	. Temperature of product	I.22. Number of packages				
		Ambient Chiled Chiled	Frozen				
	1.23	3. Identification of container/seal number	I.24. Type of packaging				
	1.25	5. Commodities certified for: Human consumption	-				
	1.26	5.	I.27. For import or admission into EU				
	1.28	3. Identification of the commodities					
	(\$	Scientific name) commodity type	roval number establishments Number Net of packages weight				
		Abatto	ir Cutting plant Cold store				

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model SUF

	II.	Health	information	II.a. Certificate reference number	II.b.			
	II.1.	Public Health Attestation						
Ę		(EC) N animal	lo 852/2004, (EC) N	sterinarian declare that I am aware of the relevar o 853/2004 and (EC) No 854/2004 and hereby uidae, Tayassuidae, or Tapiridae families describ cular that:	certify that the meat of farmed non-domestic			
Part II: Certification		II.1.1	II.1.1 the meat comes from (an) establishment(s) implementing a programme based on the HACCP principles accordance with Regulation (EC) No 852/2004;					
art II: Ce		II.1.2	 II.1.2 the meat has been obtained in compliance with the conditions set out in Section III of Annex III to Regulation (EC) No 853/2004; II.1.3 the meat fulfils the requirements of Regulation (EC) No 2075/2005 laying down specific rules on official controls for Trichinella in meat, and in particular, has been subject to an examination by a digestion method with negative results; 					
ď		II.1.3						
		II.1.4 the meat has been found fit for human consumption following ante and post-mortem inspections carried ou accordance with, Chapter II of Section I and, Chapters VII and IX of Section IV of Annex I to Regulation (I No 854/2004;						
		II.1.5		carcass or parts of the carcass have been mapter III of Section I, of Annex I to Regulation (EC				
				packages of meat have been marked with an ide lex II to Regulation (EC) No 853/2004;]	entification mark in accordance with Section I of			
		II.1.6	the meat satisfies foodstuffs;	he relevant criteria set out in Regulation (EC)	No 2073/2005 on microbiological criteria for			
II.1.7 the guarantees covering live animals and products thereof with Directive 96/23/EC, and in particular Article 29 thereof,								
II.1.8 the meat has been stored and transported in acco Regulation (EC) No 853/2004.					elevant requirements of Section I of Annex III to			
	II.2.	Anima	l Health attestation					
		I, the u	ndersigned official v	terinarian, hereby certify, that the fresh meat des	scribed in Part I:			
		II.2.1	has been obtained	n the territory/ies with code:(2)	which, at the date of issuing this certificate:			
			(¹) either [(a	has been free for 12 months from foot-and-m classical swine fever, swine vesicular disease, a				
			(¹) <i>or</i> [(a)	(i) has been free for 12 months from rinderpest, [classical swine fever] (1) and [swine vesicula				
[swine vesicu had cases/o								
 (b) during the last 12 months no vaccination against the imports of domestic animals vaccinated against the territory; 								
		11.2.2	has been obtained	rom animals that:				
				ve remained in the territory described under poinths before slaughter;]	nt II.2.1 since birth, or for at least the last three			

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model SUF

II.	Health information			II.a. Certificate reference number	II.b.
	point II.2.		point II.2	en introduced on	
	II.2.3	has been obtai	ned from	animals coming from holdings:	
		(a) in which r point II.2.1		the animals present therein have been vaccion	nated against the diseases referred to in
		(b) in and around which point II.2.1 during the		n in an area of 10 km radius, there has been no see previous 40 days,	case/outbreak of the diseases referred to in
			holdings	erinary inspections are carried out to diagnose d s are not subject to prohibition as a result of an	
	II.2.4	has been obtai	ned from	animals which:	
		(¹) either	to a	re been transported from their holdings in vehicl in approved slaughterhouse without contact with iditions mentioned above,	
	S			he slaughterhouse, have passed ante-mortem h ughter and, in particular, have shown no eviden I	
				re been slaughtered on(dd. /mm/yyyy) and(dd/mm/	
		(¹) or		re been slaughtered on the holding of origin, follo ponsible for the holding, who has provided a writ	
			_	in his opinion an unacceptable risk would have to their handlers by the transport of the animals	•
			_	the holding had been inspected and authorised of game,	by the competent authority for the slaughter
			-	the animals have passed the ante-mortem he the slaughter and, in particular, have shown point II.2.1,	
			_	the animals were slaughtered between(dd/mm/yyyy), (³)	(dd/mm/yyyy) and
			_	the bleeding of the animals was performed corn	rectly, and
			_	the slaughtered animals were eviscerated within	in three hours of the time of slaughter, and
			cor	ir carcasses have been transported to the a ditions and, where more than one hour of operature of between 0 °C and + 4 °C has been the transport;]	elapsed since the time of slaughter, a
	II.2.5	has been obtai	ned from	animals that have remained separate since birt	h from wild cloven-hoofed animals;
	II.2.6	of the disease preparation of	s referre meat for	n establishment around which, within a radius of to in point II.2.1 during the previous 40 days importation into the Union has been authorised the total cleaning and disinfection of the establishment.	or, in the event of a case of disease, the donly after slaughter of all animals present,
	II.2.7	has been obtai certificate.	ned and p	prepared without contact with other meats not co	emplying with the requirements set out in this

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model SUF

II.	Health information	II.a. Certificate reference number	II.b.
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▶⁽¹⁾ II.3. Animal welfare attestation

I, the undersigned official veterinarian, hereby certify, that the fresh meat described in Part I of this certificate derives from animals which have been handled in the slaughterhouse before and at the time of slaughter or killing in accordance with the relevant provisions of Union legislation and have met requirements at least equivalent to those laid down in Chapters II and III of Council Regulation (EC) No 1099/2009 (⁴). ◀

Notes

This certificate is meant for fresh meat, excluding offal and minced meat, of wild animals belonging to the Suidae, Tayassuidae, or Tapiridae families that are domestically kept or bred since birth in farms.

Fresh meat means all animal parts fit for human consumption, whether fresh, chilled or frozen.

Part I:

- Box reference I.8: Provide the code of territory as appearing in Part 1 of Annex II to Regulation (EU) No 206/2010.
- Box reference I.11: Place of origin: name and address of the dispatch establishment.
- Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor must inform the BIP of entry into the Union.
- Box reference I.19: Use the appropriate HS code: 02.03, 02.08.90 or 05.04.
- Box reference I.20: Indicate total gross weight and total net weight.
- Box reference I.23: For containers or boxes, the container number and the seal number (if applicable) should be included.
- Box reference I.28: Nature of commodity: Indicate 'carcass-whole', 'carcass-side', 'carcass-guarters' or 'cuts'.
- Box reference I.28: Treatment type: If appropriate indicate deboned, or bone-in. If frozen, indicate the date of freezing (mm/yy) of

Part II:

- (1) Keep as appropriate
- (2) Code of the territory as it appears in Part 1 of Annex II to Regulation (EU) No 206/2010.
- (9) Date or dates of slaughter. Imports of this meat shall not be allowed when obtained from animals slaughtered either prior to the date of authorisation for importation into the Union of the third country, territory or part thereof referred to in boxes 1.7 and 1.8, or during a period where restrictive measures have been adopted by the Union against imports of this meat from this third country, territory or

▶ ⁽²⁾	(4)	OJ L 303,	18.11.2009,	p.	1.	4
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▶ ⁽²⁾ (4) OJ L 303, 18.11.2009, p. 1. ◀					
Officia	l veterinarian				
	Name (in capital letters):	Qualification and title:			
	Date:	Signature:			
	Stamp:				

Status: Point in time view as at 01/07/2016. **Changes to legislation:** There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

Model SUW

	CO	UNTRY				veterinary certificate to E	
	1.1.	Consignor	I.2. Certificate	reference r	umber	I.2.a.	
		Name	I.3. Central Competent Authority				
		Address	I.4. Local Com				
ent		Tel. No	1.4. Local Coll	ipeterit Auti	iority		
gnm	1.5.	Consignee	I.6.				
onsi		Name					
o pe		Address					
atch		Postal code					
lispa		Tel. No					
Part I: Details of dispatched consignment	1.7.	Country ISO I.8. Region Code of origin of origin	I.9. Country of destination		de I.1	10. Region of Code destination	
Det	1.11.	. Place of origin	I.12.				
art I:		Name Approval number Address					
ď		Address					
	1.13	. Place of loading	I.14. Date of de	parture			
	I.15	. Means of transport	I.16. Entry BIP in EU				
		Aeroplane Ship Railway wagon					
		Road vehicle Other					
		Identification:	I.17.				
		Documentary references:					
	I.18	. Description of commodity	1.	l.19. Commo	dity code	e (HS code)	
					I.20. Qu	antity	
	1.21	. Temperature of product			I.22. Nu	mber of packages	
		Ambient Chiled Chiled	Frozen				
	1.23	B. Identification of container/seal number			1.24. Тур	pe of packaging	
	I.25. Commodities certified for:						
	Human consumption						
	1.26	5.	I.27. For import or admission into EU				
	1.28	3. Identification of the commodities					
	(\$	Scientific name) commodity type	roval number esta		0	Number Net of packages weight	
		Abatto	ir Cutting plar	nt Cold s	tore		

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model SUW

	II.	Health	information		II.a. Certificate reference number	II.b.		
	II.1.	Public Health Attestation						
		(EC) N the Su	lo 852/2004,(E	C) No 853/2	arian declare that I am aware of the relevant req 2004 and (EC) No 854/2004 and hereby certifidae families described in Part I was produce	y that the meat of wild animals belonging to		
Part II: Certification		II.1.1			an) establishment(s) implementing a progration (EC) No 852/2004;	amme based on the HACCP principles in		
E Cer		II.1.2	the meat has particular:	been obta	ined in accordance with Section IV of Annex	x III to Regulation (EC) No 853/2004, an in		
g.			(i) before sk	inning, it ha	s been stored and handled separately from ot	her food and not frozen;		
			and					
			(ii) after skin	ning, it has	undergone a final inspection as referred to in p	point II.1.4;		
		II.1.3 the meat fulfils the requirements of Regulation (EC) No 2075/2005 laying down specific rules on official co for <i>Trichinella</i> in meat, and in particular, has been subject to an examination by a digestion method with neg results;						
		II.1.4			if it for human consumption following a post-nil and Chapters VIII and IX of Section IV of An			
		II.1.5	(¹) either		cass or parts of the carcass have been mar			
			(¹) or		kages of meat have been marked with an ident to Regulation (EC) No 853/2004;]	ification mark in accordance with Section I of		
		II.1.6	the meat sati	meat satisfies the relevant criteria set out in Regulation (EC) No 2073/2005 on microbiological criteria for dstuffs;				
		II.1.7			live animals and products thereof provided by and in particular Article 29 thereof, are fulfilled			
		II.1.8 the meat has been stored and transported in accordance with the relevant requirements of Section I of Annex III is Regulation (EC) No 853/2004						
	II.2.		I Health attest		avian bayahu saytifu that the freeh most descri	ihad in Dart I.		
					arian, hereby certify, that the fresh meat descri			
		II.2.1			territory/ies with code:(2) which, a			
			(¹) either		been free for 12 months from foot-and-mou sical swine fever, swine vesicular disease, and			
			(¹) or		nas been free for 12 months from rinderpest, Afri classical swine fever] (1) and [swine vesicular o			
]	nas been considered free from [foot-and-mou swine vesicular disease] ('), since cases/outbreaks afterwards, and authorised to EU) No/, of((dd/mm/yyyy), without having had export this meat by Commission Regulation		
					ng the last 12 months no vaccination against orts of domestic animals vaccinated against ory;			

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model SUW

II.	Health	information		II.a. Certificate reference numbe	er	II.b.		
	II.2.2	has been obtained from wild animals that were killed between						
		, ,		eeds 20 km from the borders of a conis fresh meat into the Union,	ountry or pa	rt thereof, w	nich is not authorised du	ıring this
		(b) in an area point II.2.1;		uring the last 60 days, there has	s been no	restrictions	for the diseases referre	ed to in
	II.2.3.A	has been obtained from animals which after killing were transported within 12 hours for chilling [to a collectic centre, and immediately afterwards] (') to an approved game-handling establishment around which, within a radi of 10 km, there has been no case/outbreak of the diseases referred to in point II.2.1 during the previous 40 days in the event of a case of disease, the preparation of meat for importation into the Union has been authorised or after removal of all meat, and the total cleaning and disinfection of the establishment under the control of an officiveterinarian;				a radius days or, sed only		
(1) (4) [II.2.3.B	has been obtain negative results		arcasses on which the following te	st for classi	cal swine fev	er was carried out and p	provided
		(1) either	[virus iso	lation from blood (EDTA);]				
		(¹) or [virus isolation from samples of					;]	
		(¹) or	[immuno	fluorescence for viral antigen on sa	amples of			;]]
	II.2.4	has been obtain certificate.	ned and p	repared without contact with other	meats not c	omplying wi	th the conditions require	d in this

Notes

This certificate is meant for fresh meat, excluding offal and minced meat, of wild animals belonging to the Suidae, Tayassuidae, or Tapiridae families that are killed or hunted in the wild.

Fresh meat means all animal parts fit for human consumption whether fresh, chilled or frozen.

After importation, unskinned carcasses must be conveyed without delay to the processing establishment of destination.

Part I:

- Box reference I.8: Provide the code of territory as appearing in Part 1 of Annex II to Regulation (EU) No 206/2010.
- Box reference I.11: Place of origin: name and address of the dispatch establishment.
- Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor must inform the BIP of entry into the Union.
- Box reference I.19: Use the appropriate HS code: 02.03, 02.08.90 or 05.04.
- Box reference I.20: Indicate total gross weight and total net weight.
- Box reference I.23: For containers or boxes, the container number and the seal number (if applicable) should be included.
- Box reference I.28: Nature of commodity: Indicate 'carcass-whole', 'carcass-side', 'carcass-quarters' or 'cuts'.
- Box reference I.28: Treatment type: If appropriate, indicate 'matured' or 'unskinned'. If frozen, indicate the date of freezing (mm/yy) of the cuts/pieces.
- Box reference I.28: Abattoir: any abattoir or game handling establishment.

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model SUW

II. Health information II.a. Certificate reference number II.b.			II.b.
Par	rt II:	'	
(1)	Keep as appropriate.		
		rt 1 of Annex II to Regulation (EU) No 206/2010).
(3)	for importation into the Union of the third	uthorised when obtained from animals killed or h I country, territory or part thereof referred to in boad adopted by the Union against imports of this n	oxes reference I.7 and I.8, or during a period
(4)	with the entry 'C'. For such purpose, in	ded when required in column 5 'SG' of Part 1 o tests other than EDTA, the samples to be use uple of at least one of the following lymph nod indicated.	d are a sample of tonsil and of spleen plus
Off	icial veterinarian		
	Name (in capital letters):	Qualification	and title:
	Date:	Signature:	
		C.g. M.C.O.	
	Stamp:		

Status: Point in time view as at 01/07/2016. **Changes to legislation:** There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

Model EQW

	СО	UNTRY	Veterinary certificate to E			
	1.1.	Consignor	I.2. Certificate reference number I.2.a.			
		Name	I.3. Central Competent Authority			
		Address	I.4. Local Competent Authority			
Jent		Tel. No				
ign	1.5.	Consignee	1.6.			
suo		Name				
o pa		Address				
atch		Postal code				
disp		Tel. No	100 140 80 100 0			
Part I: Details of dispatched consignment	1.7.	Country ISO I.8. Region Code of origin code	I.9. Country of ISO I.10. Region of Code destination code destination			
: Det	1.11	. Place of origin	1.12.			
art		Name Approval number Address				
		7.00.000				
	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				
		Identification: Documentary references:	1.17.			
	I.18	. Description of commodity	I.19. Commodity code (HS code)			
			I.20. Quantity			
	1.21	. Temperature of product	I.22. Number of packages			
		Ambient Chiled Chiled	Frozen			
	1.23	B. Identification of container/seal number	I.24. Type of packaging			
	1.25					
		Human consumption				
	1.26	5.	I.27. For import or admission into EU			
	1.28	B. Identification of the commodities				
	,		umber establishments Number Net			
	(-	Scientific name) commodity Abattoir C	of packages weight Cutting plant Cold store			

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model EQW

II. Health information II.a. Certificate reference number II.b. II.1. **Public Health Attestation** I, the undersigned official veterinarian, declare that I am aware of the relevant requirements of Regulations (EC) No 178/2002, (EC) No 852/2004, (EC) No 853/2004 and (EC) No 854/2004 and hereby certify that the meat of wild solipeds belonging to the subgenus Hippotigris (zebra) described in Part I was produced in accordance with those requirements, in particular that: Part II: Certification the meat comes from (an) establishment(s) implementing a programme based on the HACCP principles in accordance with Regulation (EC) No 852/2004; II.1.2 the meat was obtained in compliance with Section IV of Annex III to Regulation (EC) No 853/2004: II.1.3 the meat fulfils the requirements of Regulation (EC) No 2075/2005 laying down specific rules on official controls for Trichinella in meat, in particular, has been subject to an examination by a digestion method with negative results; the meat has been found fit for human consumption following a post-mortem inspection carried out in accordance with Chapter II of Section I and Chapters VIII and IX of Section IV of Annex I to Regulation (EC) No 854/2004: II.1.5 (1) either [the carcass or parts of the carcass have been marked with a health mark in accordance with Chapter III of Section I of Annex I to Regulation (EC) No 854/2004;] Ithe packages of meat have been marked with an identification mark in accordance with Section I of (1) or Annex II to Regulation (EC) No 853/2004;] II.1.6 the meat satisfies the relevant criteria set out in Regulation (EC) No 2073/2005 on microbiological criteria for foodstuffs: II.1.7 the guarantees covering live animals and products thereof provided by the residue plans submitted in accordance with Directive 96/23/EC, and in particular Article 29 thereof, are fulfilled; II.1.8 the meat has been stored and transported in accordance with the relevant requirements of Section I of Annex III to Regulation (EC) No 853/2004. 11.2. **Animal Health attestation** I, the undersigned official veterinarian, hereby certify, that the fresh meat described in Part I: (dd/mm/yyyy) and II.2.1 has been obtained from wild animals that were killed between has been obtained from wild animals which after killing were transported within 12 hours for chilling [to a collection centre, and immediately afterwards] (1) to an approved game-handling establishment around which, within a radius of 10 km, there has been no case/outbreak of African horse sickness or glanders during the previous 40 days or, in the event of a case of such diseases, the preparation of meat for exportation to the Union has been authorised only after removal of all meat, and the total cleaning and disinfection of the establishment under the control of an official II.2.3 has been obtained and prepared without contact with other meats not complying with the requirements set out in this certificate. Notes This certificate is meant for fresh meat, excluding offal and minced meat, of wild solipeds belonging to the subgenus Hippotigris (zebra). Fresh meat means all animal parts fit for human consumption whether fresh, chilled or frozen.

After importation, unskinned carcasses must be conveved without delay to the processing establishment of destination.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model EQW

II.	Health information	II.a. Certificate reference number	II.b.
Part — E — E — E — E — E — E — E — E — E —	I: Box reference I.8: Provide the code of to Box reference I.11: Place of origin: name Box reference I.15: Registration numbe provided. In case of unloading and reloadox reference I.19: Use the appropriate Box reference I.20: Indicate total gross of Box reference I.23: For containers or both Box reference I.28: Nature of commodity Box reference I.28: Nature of commodity Box reference I.28: Nature of commodity Box reference I.28: Abattoir: any abattoid II: Keep as appropriate. Dates. Imports of this meat shall not be an or importation into the Union of the thin estrictive measures have been adopted.	erritory as appearing in Part 1 of Annex II e and address of the dispatch establishr r (railway wagons or container and lorrie ding, the consignor must inform the BIF HS code: 02.08.90 or 05.04. weight and total net weight. xes, the container number and the seal of the container number and the seal of the container number and the seal of the container or unskinner or game handling establishment. uthorised when obtained from animals kill discountry, territory or part thereof referred	to Regulation (EU) No 206/2010. lent. s), flight number (aircraft) or name (ship) is to be of entry into the Union. lumber (if applicable) should be included. ', 'carcass-quarters' or 'cuts'. 'd'. If frozen, indicate the date of freezing (mm/yy) ed or hunted either prior to the date of authorisation d to in boxes 1.7 and 1.8, or during a period where at from this third country, territory or part thereof.
Offici	ial veterinarian		
	Name (in capital letters):	Qualif	cation and title:
	Date:	Signat	ure:
	Stamp:		

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

Model NZ-TRANSIT-SG

col	JNTR	<i>r</i> :	Veterina	ary certificate to EU		
	l.1.	Consignor	I.2. Certificate reference number	I.2.a.		
		Name	I.3. Central Competent Authority			
		Address	I.4. Local Competent authority			
		Country				
_		Tel.				
men	1.5.	Consignee	1.6.			
sign		Name				
COU		Address				
hed						
patc		Country Tel.				
gip	1.7.	Country ISO I.8. Region Code	I.9. Country of ISO code I.10	<u> </u>		
ls of	1.7.	of origin code of origin	destination	J.		
Detai	Si	ngapore SG				
Part I: Details of dispatched consignment	1.11.	Place of origin	I.12.			
Pai						
		Name Approval number				
		Address				
	112	Place of leading	114 Data of departure Time	o of donortura		
	1.13.	Place of loading Address	I.14. Date of departure Time	e 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 CITES			
		Identification:				
		Document:				
	I.18.	Description of commodity	I.19. Commodity	code (HS code)		
				D. Quantity		
	1.21.	Temperature of product	1.22	Number of packages		
		Ambient ☐ Chilled ☐	Frozen 🗆	paramage of		
	123	Seal/Container No		4. Type of		
	1.20.	Cean Container 140	1.2-	packaging		
	1.25.	Commodities certified as:				
		Human consumption				
	1.26.		I.27. For import or admission into EU	Ц		
		Identification of the commodity				
		Species Nature of commodity Appro- scientific	oval number of establishments No	umber Net weight of		
		name) Abattoir	Cutting plant Cold store page	ckages		

Part II: Certification

Document Generated: 2024-06-03

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model NZ-TRANSIT-SG

II. Health information II.a. Certificate reference number II.b.

II.1 Health attestation

- I, the undersigned official veterinarian, hereby certify, that the fresh meat described in Part I:
- II.1.1 originates from New Zealand and is authorised for introduction into the Union as laid down in Part 1 of Annex II to Regulation (EU) No 206/2010, and
- II.1.2 is destined for the Union and is accompanied by the veterinary certificate drawn up in accordance with the model set out in Annex I to Commission Implementing Decision (EU) 2015/1901 (¹) issued by the competent authority of New Zealand with certificate reference number, and
- II.1.3 during transit has been unloaded, stored, reloaded and transported in accordance with the relevant
 ▶ "requirements of Section I and V respectively of Annex III to Regulation (EC) No 853/2004 ◄, and
- II.1.4 during all stages of transit has been kept segregated from animal products not eligible for import into the Union, and
- II.1.5 is eligible for import into the Union.

II.2 Transit attestation

- I, the undersigned official veterinarian, hereby certify, that the consignment of fresh meat described in Part I has:
- II.2.1 arrived to the customs area of Singapore airport, in cartons with at least one tamper proof seal applied on outer packaging of each carton in such a way, that the cartons cannot be opened without at least one seal is destroyed or damaged, and
- II.2.2 immediately after unloading from the plane, been subject to documentary and identity check and if applicable physical check (2) by the competent authority of Singapore, and
- II.2.3 been stored in an approved establishment in the customs area of Singapore (3), and
- II.2.4 been reloaded into a reefer container in an approved establishment in the customs area of Singapore under supervision of the competent authority of Singapore, and

the reefer container has been:

- II.2.5 sealed by the Customs authority of Singapore, for transport from the approved establishment to the sea port of Singapore, and
- II.2.6 sealed by the competent authority of Singapore, for transport from the approved establishment until arrival at the first Union border inspection post.

Notes

This certificate is meant for the following commodities of fresh meat originating from New Zealand and for which New Zealand is authorised to introduce into the Union, which is accompanied by the appropriate model of veterinary certificate issued by the competent authority of New Zealand, destined to the Union and being unloaded, reloaded and transited with or without storage through Singapore:

- fresh meat, including minced meat, of:
 - (1) domestic bovine animals (including *Bubalus* and Bison species and their cross-breeds);
 - (2) domestic ovine animals (Ovis aries) or domestic caprine animals (Capra hircus);
 - (3) domestic porcine animals (Sus scrofa);
 - (4) domestic solipeds (Equus caballus, Equus asinus and their cross-breeds);

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUN	COUNTRY			Model NZ-TRANSIT-S		
II.	Health	information	II.a.	Certificate reference number	II.b.	
_	fresh m	eat, excluding offal and minced r	neat, c	of:		
	(5)		cross-	order <i>Artiodactyla</i> (excluding bovine anima breeds), <i>Ovis aries, Capra hircus</i> , <i>Suidae</i> d <i>Elephantidae</i> ;		
	(6) wild non-domestic animals of the order Artiodactyla (excluding bovine animals (including Bison and Bubalus species and their cross-breeds), Ovis aries, Capra hircus, Suidae and Tayassuidae), and of the families Rhinocerotidae and Elephantidae;					
	(7)	farmed non-domestic animals b	elongi	ng to the <i>Suidae, Tayassuidae</i> , or <i>Tapirida</i>	e families;	
	(8)	wild non-domestic animals belo	nging	to the <i>Suidae, Tayassuidae</i> , or <i>Tapiridae</i> fa	amilies.	
	Fresh n	neat means all animal parts fit for	huma	n consumption whether fresh, chilled or fro	ozen.	
Part	l:					
_	Box ref	erence I.7: Country of origin mea	ns her	e the country of dispatch: Singapore.		
_	Box ref Singap		ne, ad	dress and approval number of the dispato	h establishment in	
_	Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor must inform the BIP of entry into the Union.					
_		ference I.19: Use the appropria 05.04 or 15.02.	te HS	code: 02.01, 02.02, 02.03, 02.04, 02.05	, 02.06, 02.08.90,	
_	Box ref	erence I.20: Indicate total gross v	veight	and total net weight.		
_		erence I.23: For containers: The ent authority of Singapore at the		iner number and the seal number of the setion of reloading.	seal applied by the	
_				cate 'carcass-whole', 'carcass-side', 'carca ne approved establishments in New Zealan		
Part	II:					
(1)	For consignments of fresh meat for which equivalence has been determined under the Agreement between the European Community and New Zealand (Council Decision 97/132/EC), the appropriate model veterinary certificate is set out in Annex I to Commission Implementing Decision (EU) 2015/1901 of 20 October 2015 laying down certification rules and a model health certificate for importation into the Union of consignments of live animals and animal products from New Zealand and repealing Decision 2003/56/EC.				appropriate model EU) 2015/1901 of ation into the Union	
(²)	In exce	eptional cases which may prese ted, additional physical checks m	nt a p ust be	ublic health or animal health risk or whe carried out.	n irregularities are	
(³)	Delete	if the consignment has been relo	aded v	vithout storage.		
Offic	ial veterii	narian				
	Name (in capital letters):		Qualification and tit	le:	
	Date:			Signature:		
	Stamp:					

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

Textual Amendments

F20 Substituted by Commission Regulation (EU) No 810/2010 of 15 September 2010 amending Regulation (EU) No 206/2010 laying down lists of third countries, territories or parts thereof authorised for the introduction into the European Union of certain animals and fresh meat and the veterinary certification requirements (Text with EEA relevance).

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

ANNEX III

Model TRANSIT/STORAGE

	CO	UNTRY	Veterinary certificate to EU
	1.1.	Consignor	I.2. Certificate reference number I.2.a.
		Name	I.3. Central Competent Authority
		Address	
ent		Tel. No	I.4. Local Competent Authority
gnm	1.5.	Consignee	I.6. Person responsible for the consignment in EU
onsi		Name	Name
o pe		Address	Address
tche		Postal code	Postal code
ispa		Tel. No	Tel. No
Part I: Details of dispatched consignment	1.7.	Country ISO I.8. Region Code of origin	I.9. Country of ISO I.10. Region of Code destination code destination
Deta	1.11.	Place of origin	I.12. Place of destination
#		Name Approval number	Custom warehouse Ship supplier
Pa		Address	Name Approval number
			Address Postal code
	I.13	. Place of loading	I.14. Date of departure
	1.15	. Means of transport	I.16. Entry BIP in EU
		Aeroplane Ship Railway wagon	
		Road vehicle Other	
		Identification:	I.17. No. (s) of CITES
		Documentary references:	
	I.18	. Description of commodity	I.19. Commodity code (HS code)
			I.20. Quantity
	1.21	. Temperature of product	I.22. Number of packages
		Ambient Chiled	Frozen
	1.23	. Identification of container/seal number	I.24. Type of packaging
	1.25	. Commodities certified for:	·
		Human consumption	
	1.26	. For transit through EU to 3 rd Country	1.27.
		3rd country ISO code	
	1.28	. Identification of the commodities	
	(\$	Species Nature of Treatment Approval nu Scientific name) commodity type	umber establishments Number Net of packages weight
		Abattoir	Cutting manufacturing plant/ plant

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY	Model TRANSIT/STORAGE

	II.	Health i	nformation	II.a. Certificate reference number	II.b.				
	II.1.	Animal	Health Attestation						
		I, the undersigned official veterinarian, hereby certify, that the fresh meat described in Part I:							
_		II.1.1	comes from a country or (EU) No 206/2010 at the	region authorized for imports into the Union as time of slaughter, and	laid down in Part 1 of Annex II to Regulation				
Part II: Certification				nt animal health conditions as laid down in OR] [EQU] [RUF] [RUW] [SUF] [SUW] [EQW					
Part II: C		II.1.3		which were slaughtered and processed on (dd/mm/yyyy) and					
	Notes								
				ge in accordance with Article 12(4) or Article 1	3 of Directive 97/78/EC of:				
	— fresi		cluding minced meat, of:						
	(1)			ng <i>Bubalus</i> and <i>Bison</i> species and their cross					
	(2)			es) or domestic caprine animals (Capra hircus) (Model 'OVI');				
	(3)		ic porcine animals (Sus so	crofa) (Model 'POR');					
	_ fresi		cluding minced meat, of:						
	(4)			us, Equus asinus and their cross-breeds) (Mod	del 'EQU');				
	— fresi	h meat, ex	cluding offal and minced	meat, of:					
	(5)		ss-breeds), Ovis aries, Ca	the order Artiodactyla (excluding bovine anima apra hircus, Suidae and Tayassuidae), and of th					
	(6)		ss-breeds), Ovis aries, Ca	e order Artiodactyla (excluding bovine animals apra hircus, Suidae and Tayassuidae), and of th					
	(7)	farmed	non-domestic animals be	longing to the Suidae, Tayassuidae, or Tapirida	ae families (Model 'SUF');				

wild non-domestic animals belonging to the Suidae, Tayassuidae, or Tapiridae families (Model 'SUW');

wild solipeds belonging to the subgenus Hippotigris (zebra) (Model 'EQW'). Fresh meat means all animal parts fit for human consumption whether fresh, chilled or frozen.

(9)

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model TRANSIT/STORAGE

II.	Health information	II.a. Certificate reference number	II.b.				
Pa	Part I:						
	Box reference I.8: Provide the code of the Box reference I.11: Place of origin: name Box reference I.12: Address (and approor of ship chandler shall be included. Box reference I.15: Registration number provided. In case of unloading and reloaded. Box reference I.19: Use the appropriate Box reference I.20: Indicate total gross of Box reference I.23: For containers or both Box reference I.28: Nature of commoditing Box reference I.28: Treatment type: If from the III: Keep as appropriate. Date or dates of slaughter. Imports of the date of authorisation for exportation to the same properties.	erritory as appearing in Part 1 of Annex II to Re e and address of the dispatch establishment. val number if known) of the warehouse in a fre r (railway wagons or container and lorries), fliading, the consignor must inform the BIP of er HS code: 02.01, 02.02, 02.03, 02.04, 02.05, weight and total net weight. exes, the container number and the seal numb y: Indicate 'carcass-whole', 'carcass-side', 'carcase, indicate the date of freezing (mm/yy) of the seal of the third country, territory or part the verbeen adopted by the Union against imports	ght number (aircraft) or name (ship) is to be try into the Union. 02.06, 02.08.90, 02.09, 05.04 or 15.02. er (if applicable) should be included. ccass-quarters', 'cuts', or 'minced meat'. the cuts/pieces.				
Off	iicial veterinarian						
	Name (in capital letters):	Qualification	n and title:				
	Date:	Signature:					
	Stamp:						

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

ANNEX IV

ANIMALS REFERRED TO IN ARTICLE 1(1)(b)

PART 1

Lists of third countries, territories or parts thereof

SECTION 1

Parts of third countries or territories referred to in Article 7(2)

[F20Country/territory	Code of part of the country/territory	Description of part of the country/territory
US – United States	US-A	The State of Hawaii ^a
a Suspended from 5 May 2010.]		

PART 2

Tables of animals and the corresponding model veterinary certificates

Table 1				
'QUE'	: Model of veterinary certificate for consignments of queen bees and queen bumble bees (<i>Apis mellifera and Bombus</i> spp.),			
'BEE'	: Model of veterinary certificate for consignments of colonies of bumble bees (<i>Bombus</i> spp.)			
Order	Family	Genera/species		
Hymenoptera	Apidae	Apis mellifera, Bombus spp.		

[F21Model QUE]

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

cou	INTR	1					Veterinary cert	ificate to EU
	l.1.	Consignor Name			reference		1.2.a.	
	Address		1.3.	Central c	ompetent a	uthority		
ıı		Tel.	1.4.	Local cor	npetent au	thority		
l me	1.5.	Consignee	1.6.					
ısig		Name						
8		Address						
hed		Postal code						
dispatched consignment		Tel.						
5	1.7.	Country of origin ISO code I.8. Region of origin Code	1.9.	Country of destination		O code	I.10. Region of destination	Code
Detai	l.11.	Place of origin	I.12.	Place of	destination			
Part I: Details		Name Approval number Address						
	I.13.	Place of loading	1.14.	Date of d	eparture			
	Address Approval number							
	l.15.	Means of transport	I.16. Entry BIP in EU					
		Aeroplane ☐ Ship ☐ Railway wagon ☐						
		Road vehicle Other						
		Identification	1.17.	No(s) of	CITES			
		Documentary references						
	I.18.	Description of commodity	I.19. Commodity code (HS code)					
						01.0	6.41	
						1.2	0. Quantity	
	1.21.					1.2	2. Number of packages	;
	1.23.	Identification of container/seal number				1.2	4.	
	1.25.	Commodities certified for:						
		Breeding □						
	1.26.		1.27.	For impor	t or admis	sion into	EU 🗆	
	1.28.	Identification of the commodities						
		Species (scientific name)						
	1							

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

	COUNTRY						
	II.	Health information	II.a. Certificate reference number	II.b.			
	II.1.	Animal Health attestation					
	I, the undersigned, hereby certify, that the animals referred to in Part I of this certificate meet the following requirements:						
5	II.1.1.	I.1.1. they come from the territory with code:					
rtificati	II.1.2.	they:					
II.1.2. they: (a) come from a breeding apiary, which is supervised and controlled by the competent authority;							
		(b) come from an area which is not subject to any restrictions a occurrence has taken place within at least 30 days prior to foulbrood has occurred previously, all hives within a radius o infected hives burned or treated and inspected to the satist recorded case:	the issuance of the present certificat f three kilometres have been checked	e. Where an outbreak of American by the competent authority and all			
		(c) are from hives or come from hives or colonies (in the case o last 30 days for American foulbrood as laid down in the O negative results;					
		(d) come from an area of at least 100 km radius which is not subeetle (Aethina tumida) or Tropilaelaps spp., and where the		ith the occurrence of the small hive			
		(e) are from hives or come from hives or colonies (in the case of show no clinical signs or suspicion of disease including infer		d immediately prior to dispatch and			
		(f) Have undergone detailed examinations to ensure that all bee their eggs and larvae, or other infestations, in particular <i>Trop</i>		mall hive beetle (Aethina tumida) or			
	II.1.3.	the packaging material, queen cages, accompanying products brood-combs, and all precautions have been taken to prevent companying products and the packaging material, queen cages, accompanying products are companying products.					
	Notes						
	Part I:						
	Mer	reference I.12: the introduction of queen bees and their accompnder States listed in the third column of the table set out in the 0.2013, p. 38).					
	— Box reference I.20: Number of queen bees (Apis mellifera and Bombus spp.). Each queen bee may be accompanied by a maximum of 20 attendants.						
	Part II:						
	(1) Code of the territory as it appears in Part 1 of Annex II or Section 1 of Part 1 of Annex IV to Commission Regulation (EU) No 206/2010.						
	Official	veterinarian/Official inspector					
	Na	me (in capital letters):	Qualifica	tion and title:			
	Da	te:	Signature	э:			
	Sta	amp:					

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

Model BEE

	CO	UNTRY	Veterinary certificate to EU
	1.1.	Consignor	I.2. Certificate reference number I.2.a.
		Name	I.3. Central Competent Authority
		Address	
		Tel. No	I.4. Local Competent Authority
ıt	1.5.	Consignee	1.6.
mer		Name	
sign		Address	
con		Postal code	
hed		Tel. No	
Part I: Details of dispatched consignment	1.7.	Country ISO I.8. Region Code of origin code	I.9. Country of ISO I.10. Region of Code destination code destination
of d			
tails	1.11.	. Place of origin	1.12.
Det		Name Approval number Address	
Part		Name Approval number Address	
		Name Approval number Address	
	1.13	. Place of loading	I.14. Date of departure time of departure
		Address Approval number	
	1.15	. Means of transport	I.16. Entry BIP in EU
		Aeroplane Ship Railway wagon	
		Road vehicle Other	I.17. No(s) of CITES
		Identification: Documentary references:	
	I.18	. Description of commodity	I.19. Commodity code (HS code) 01.06.90
			I.20. Quantity
	1.21		I.22. Number of packages
	1.23	B. Identification of container/seal number	1.24.
	1.25	5. Commodities certified for:	
		Breeding	
	1.26	5.	I.27. For import or admission into EU
	1.28	B. Identification of the commodities	
			fication Identification stem number

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY	Model BE
COUNTRY	Widdel BE

	II.	Health information	II.a. Certificate reference number	II.b.			
	II.1.	Animal Health attestation:					
		I, the undersigned, hereby certify that:					
	II.1.1						
ication	 (a) the bumble bees (Bombus spp.) referred to in Part I of this certificate have been bred and kept under a controlled environment within a recognised establishment which is supervised and controlled by the competent authority; 						
Part II: Certification			eferred to in Part I of this certificate was inspeeding stock show no clinical signs or suspici				
Pa		broodstock and pac	ort into the Union have undergone detailed a kaging do not contain the small hive beetle (A ular <i>Tropilaelaps</i> spp., affecting bees;				
			ontainers, accompanying products and food combs, and all precautions have been taken to fbees.				
	Notes						
	Part I:						
		reference I.20: Number of contain- ble bees.	ers of bumble bees (<i>Bombus</i> spp.), each cor	ntaining a colony of a maximum of 200 adult			
	Official v	eterinarian /Official inspector					
		Name (in capital letters):	Qualificatio	n and title:			
		Date:	Signature:				
		Stamp:					

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

ANNEX V

Explanatory notes for completing the veterinary certificates (referred to in Article 18)

- (a) Veterinary certificates shall be issued by the exporting third country, based on the models set out in Part 2 of Annexes I, II and IV and Annex III according to the layout of the model that corresponds to the live animals/fresh meat concerned.
 - They shall contain, in the numbered order that appears in the model, the attestations that are required for any third country and, as the case may be, those supplementary guarantees that are required for the exporting third country or part thereof.
 - If the Member State of destination imposes, for the live animals/fresh meat concerned, additional certification requirements, attestations to certify that those requirements are fulfilled shall also be incorporated in the original form of the veterinary certificate.
- (b) Where the model 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 the certificate.
- (c) A separate and unique certificate must be provided for the live animals/fresh meat that are exported from a territory or territories of the same exporting country appearing in columns 2 and 3 of Part 1 of Annex I, II or IV which are consigned to the same destination and transported in the same railway wagon, lorry, aircraft or ship.
- (d) The original of each 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.
- (e) The veterinary 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 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.
- (f) If for reasons of identification of the items of the consignment (schedule in point I.28 of the model veterinary certificate), additional sheets of paper are attached to the certificate, those sheets of paper shall also be considered as forming part of the original of the certificate by the application of the signature and stamp of the certifying officer, on each of the pages.
- (g) When the certificate, including additional schedules referred to in (f), 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 at the top of the pages.
- (h) The original of the certificate must be completed and signed by an official veterinarian or by another designated official inspector where this is provided for in the model veterinary certificate. In the case of live animals, the certificate must be completed and signed within 24 hours prior to loading of the consignment for introduction into the Union. The competent authorities of the exporting third country shall ensure that rules of certification equivalent to those laid down in Directive 96/93/EC⁽¹⁰⁾ are followed.

The colour of the signature shall be different from that of the printing. This requirement also applies to stamps other than those embossed or watermarked.

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

(i) The certificate reference number referred to in boxes I.2 and II.a. must be issued by the competent authority.]

[F22ANNEX VI

Textual Amendments

F22 Inserted by Commission Implementing Regulation (EU) No 780/2013 of 14 August 2013 amending Commission Regulation (EU) No 206/2010 laying down lists of third countries, territories or parts thereof authorised for the introduction into the European Union of certain animals and fresh meat and the veterinary certification requirements (Text with EEA relevance).

PART 1

Table 1		
'RUM-A' :		for animals of the species listed below ntended for an approved body, institute
Order	Family	Genera/species
Artiodactyla	Antilocapridae	Antilocapra ssp.
	Bovidae	Addax ssp., Aepyceros ssp., Alcelaphus ssp., Ammodorcas ssp., Ammotragus ssp., Antidorcas ssp., Bos ssp. (including Bibos, Novibos, Poephagus), Boselaphus ssp., Bubalus ssp. (including anoa), Budorcas ssp., Capra ssp., Cephalophus ssp., Connochaetes ssp., Damaliscus ssp. (including Beatragus), Dorcatragus ssp., Gazella ssp., Hemitragus ssp., Hippotragus ssp., Kobus ssp., Litocranius ssp., Madoqua ssp., Naemorhedus ssp. (including Nemorhaedus and Capricornis), Neotragus ssp., Oreamnos ssp., Oreotragus ssp., Oryx ssp., Ourebia ssp., Ovibos ssp., Ovis ssp., Patholops ssp., Pelea ssp., Procapra ssp., Pseudois ssp., Pseudoryx ssp.,

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

		Raphicerus ssp., Redunca ssp., Rupicapra ssp., Saiga ssp., Sigmoceros-Alecelaphus ssp., Sylvicapra ssp., Syncerus ssp., Taurotragus ssp., Tetracerus ssp., Tragelaphus ssp. (including Boocerus).
	Camelidae	Camelus ssp., Lama ssp., Vicugna ssp.
	Cervidae	Alces ssp., Axis-Hyelaphus ssp., Blastocerus ssp., Capreolus ssp., Cervus-Rucervus ssp., Dama ssp., Elaphurus ssp., Hippocamelus ssp., Hydropotes ssp., Mazama ssp., Megamuntiacus ssp., Muntiacus ssp., Odocoileus ssp., Ozotoceros ssp., Pudu ssp., Rangifer ssp.
	Giraffidae	Giraffa ssp., Okapia ssp.
	Moschidae	Moschus ssp.
	Tragulidae	Hyemoschus ssp., Tragulus- Moschiola ssp.
Table 2		
th		or animals of the species listed below ended for an approved body, institute
Order	Family	Genera/species
Artiodactyla	Suidae	Babyrousa ssp., Hylochoerus ssp., Phacochoerus ssp., Potamochoerus ssp., Sus ssp.
	Tayassuidae	Catagonus ssp., Pecari- Tayassu ssp.
	Hippopotamidae	Hexaprotodon-Choeropsis ssp., Hippopotamus ssp.
Table 3		
'TRE-A' : M		for animals of the species listed below ended for an approved body, institute
Order	Family	Genera/species
	١ -	

Status: Point in time view as at 01/07/2016. **Changes to legislation:** There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

		Ceratotherium ssp., Dicerorhinus ssp., Diceros ssp., Rhinoceros ssp.
Proboscidea	Elephantidae	Elephas ssp., Loxodonta ssp.

PART 2 Model RUM-A

cou	INTR	1	Veterinary certificate to EU		
	l.1.	Consignor Name	I.2. Certificate reference No I.2.a.		
		Address	I.3. Central competent authority		
ent		Tel.	1.4. Local competent authority		
ignm	1.5.	Consignee Name	1.6.		
dispatched consignment		Address			
ched		Postal code			
spat		Tel.	100 111 110 2 111 110 2		
ō	1.7.	Country of origin ISO code I.8. Region of origin Code	I.9. Country of destination ISO code l.10. Region of destination Code		
Deta	l.11.	Place of origin	1.12.		
Part I: Details		Name Approval number Address			
	I.13.	Place of loading	I.14. Date of departure		
		Address Approval number	·		
	l.15.	Means of transport	I.16. Entry BIP in EU		
		Aeroplane Ship Railway wagon Railway wagon			
		Road vehicle Other Ildentification	1.17.		
		Documentary references			
	I.18.	Description of commodity	I.19. Commodity code (HS code)		
			I.20. Quantity		
	I.21.		I.22. Number of packages		
	1.23.	Seal/Container No	1.24.		
	1.25.	Commodities certified for:			
		Approved body			
	1.26.		I.27. For import or admission into EU		
	1.28.	Identification of the commodities			
		Species Identification system (scientific name)	Identification number Age Sex		

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model RUM-A

II. Health information II.a. Certificate reference number II.b.

II.1. Animal health attestation

Part II: Certification

I, the undersigned official veterinarian responsible for the approved body, institute or centre/holding (1) of origin certify that the animals described in Part I meet the following requirements:

- II.1.1. They come from the country, territory or part thereof described in Box I.7.:
 - (a) where the diseases referred to in this certificate are notifiable,
 - ▶ (1) (b) which at the date of issuing this certificate has been free for 12 months from rinderpest. ◄
- II.1.2. They come from the body, institute or centre/holding (1) described in Box I.11;
 - (a) which is approved according to the requirements and conditions set out in Part 3 and 4 of Annex VI to Regulation (EU) No 206/2010;
 - (b) which is not subjected to any restrictions relating to a national programme for the control of infectious diseases to which the animals referred to in Box I.28. are susceptible;
 - (c) where there have been no clinical cases of the following diseases to which the animals referred to in Box I.28. are susceptible:
 - anthrax for the last 30 days;
 - foot-and-mouth disease, bluetongue, Rift valley fever, vesicular stomatitis, rabies, contagious bovine pleuropneumonia, lumpy skin disease, peste des petits ruminants, sheep pox, goat pox, contagious caprine pleuropneumonia for the past 6 months;
 - (d) where there have been no clinical or non-clinical cases of tuberculosis and brucellosis for the past 6 months;
 - (e) around which in an area of 10 km radius for the last 30 days, there has been no case of the following diseases to which the animals referred to in Box I.28. are susceptible: foot-and-mouth disease, vesicular stomatitis, contagious bovine pleuropneumonia, peste des petits ruminants, sheep pox, goat pox, contagious caprine pleuropneumonia;
 - (f) around which in an area of 150 km radius for the last 30 days, there has been no case of the following diseases to which the animals referred to in Box I.28. are susceptible: bluetongue, epizootic haemorrhagic disease, Rift valley fever, lumpy skin disease;
 - (g) in which they have remained since birth or for the past 6 months before dispatch to the Union.

II.1.3. Thev:

- (a) have not come into contact with other animals not complying with at least the same health requirements as described in this certificate for the last 30 days and during their transportation from the approved body, institute or centre/holding (1) to the place of shipment;
- (b) were examined by an official veterinarian within 24 hours of loading and showed no clinical sign of disease and are fit for the intended transport;
- (c) are not animals to be killed under a national programme for the eradication of diseases.

II.1.4. Foot-and-Mouth Disease

- either (1) [(a) They come from the country, territory or part thereof described in Box I.7 which has been free for the past 12 months from foot-and-mouth disease with or without vaccination, and]
- or (1) [(a) They have been subjected to the following tests:
 - a serological test for evidence of foot-and-mouth disease virus infection carried out in accordance with one of the
 prescribed tests for international trade laid down in the OIE Manual of Diagnostic Tests and Vaccines for Terrestrial
 Animals (OIE Terrestrial Manual), with negative results, taken within 10 days prior to dispatch to the Union,
 - (1)(2)[a probang test for evidence of foot-and-mouth disease virus infection carried out in accordance with the procedures described in the OIE Terrestrial Manual with negative results, (1)(2)[taken 10 days prior to dispatch to the Union] (1)(4)[taken on two occasions 15 days apart, the second of which must have been taken 10 days prior to dispatch to the Union, and]
- ▶ (2) (1) (b) they have not been vaccinated against foot-and-mouth disease. ◄

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model RUM-A Health information II.a. Certificate reference number II b Bluetongue and Epizootic haemorrhagic disease (EHD) [They come from the country, territory or part thereof described in Box 1.7 which has been free for 24 months from blue-tongue/EHD in accordance with the OIE Terrestrial Animal Health Code (OIE Terrestrial Code).] either (1) [They were held in a vector-protected facility in the approved body, institute or centre/holding (1) for at least 30 days prior to shipment and were subjected to a serology test according to the OIE Terrestrial Manual, with negative results, carried out at least 28 days after introduction into the approved body, institute or centre.] or (1) [They were held in a vector-protected facility in the approved body, institute or centre/holding (¹) for at least 30 days prior to shipment and were subjected to a PCR test according to the OIE Terrestrial Manual, with negative results, carried out at least 14 days after introduction into the approved body, institute or centre.] or (1) or (1) [They come from a seasonally free area and were subjected during that period to an serology test according to the OIE Terrestrial Manual, with negative results, carried out at least 28 days after introduction into the approved body, institute or [They come from a seasonally free area and were subjected during that period to a PCR test according to the OIE Terrestrial Manual, with negative results, carried out at least 14 days after introduction into the approved body, institute or centre/holding (1).] II.1.6. Rift valley fever [They come from the country, territory or part thereof described in Box I.7. which has been free for 48 months from Rift valley fever and have not been vaccinated against that disease.] either (1) [They were held in a vector-protected facility in the approved body, institute or centre/holding (¹) for at least 30 days prior to shipment during which the animals showed no clinical signs of Rift valley fever and were protected from vectors between the vector-protected facility and the place of shipment to the Union as well as at the place of shipment.] or (1) [They have been subjected to a virus neutralisation test (a) with negative results for evidence of Rift valley fever, as laid down and prescribed for international trade by the OIE Terrestrial Manual, taken at the beginning of the isolation/quarantine period and at least 42 days later on, the second of which must have been taken ▶ within 10 days prior to dispatch to the Union. ■] or (1) II.1.7. Brucellosis [They come from a country, territory or part thereof described in Box I.7 which has been free for the past 12 months from brucellosis and which have not been vaccinated against that disease;] either (1) or (1) [They have been subjected to a test as laid down and prescribed for international trade by the OIE Terrestrial Manual, in the 30 days prior to dispatch to the Union:1 or (1) They are castrated males of any agel. II.1.8. Other vaccinations (a) They have not been vaccinated against vesicular stomatitis, (5) (b) They have been vaccinated against: (dd/mm/yyyy)(date(s)) with the following vaccine(s) (name of vaccine(s) med on (dd/mm/yyyy)(date(s)) shows a protective immune response.]. used) and a blood test performed on .. II.1.9. Parasite treatment They have been treated at least twice during the 40 days prior to dispatch to the Union against internal and external parasites II.1.10. Loading on the means of transport

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model RUM-A

II. Health information	II.a. Certificate reference number	II.b.
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Notes

This certificate is to be used for live animals listed in the note for Box I.28. coming from an approved body, institute or centre in a third country, territory of part thereof, and destined to an approved body, institute or centre situated within a Member State. Use one certificate per species.

Part I:

Box reference I.15.: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor shall inform the BIP of entry into the EU.

Box reference I.19.: Use appropriate HS code: 010613 or 010619.

Identification system: Specify the identification system (tag, tattoos, brand, chip, transponder). The identifier shall include the ISO code of the exporting country and permit tracing of their premises of origin. Box reference I.28.:

Age: months.

Sex (M = male, F = female, C = castrated).

Species: Select the species amongst those listed below:

Order	Family	Genera/species
Artiodactyla	Antilocapridae	Antilocapra
	Bovidae	Addax ssp., Aepyceros ssp., Alcelaphus ssp., Ammodorcas ssp., Ammotragus ssp., Antidorcas ssp., Antidope ssp., Bison ssp., Bos ssp. (including Bibos, Novibos, Poephagus), Boselaphus ssp., Bubalus ssp. (including anoa), Budorcas ssp., Capra ssp., Cephalophus ssp., Connochaetes ssp., Damaliscus ssp. (including Beatragus), Dorcatragus ssp., Gazella ssp., Hemitragus ssp., Hippotragus ssp., Kobus ssp., Litocranius ssp., Madoqua ssp., Naemorhedus ssp. (including Nemorhaedus and Capricornis), Neotragus ssp., Oreamnos ssp., Oreotragus ssp., Oryx ssp., Ourboia ssp., Ovibo ssp., Ovis ssp., Patholops ssp., Peleda ssp., Pseudois ssp., Ssp., Pseudoryx ssp., Raphicerus ssp., Redunca ssp., Rupicapra ssp., Saiga ssp., Sigmoceros-Alecelaphus ssp., Sylvicapra ssp., Syncerus ssp., Taurotragus ssp., Tetracerus ssp., Tragelaphus ssp. (including Boocerus).
	Camelidae	Camelus ssp., Lama ssp., Vicugna ssp.
	Cervidae	Alces ssp., Axis-Hyelaphus ssp., Blastocerus ssp., Capreolus ssp., Cervus-Rucervus ssp., Dama ssp., Elaphurus ssp., Hippocamelus ssp., Hydropotes ssp., Mazama ssp., Megamuntiacus ssp., Muntiacus ssp., Odocoileus ssp., Ozotoceros ssp., Pudu ssp., Rangifer ssp.
	Giraffidae	Giraffa ssp., Okapia ssp.
	Moschidae	Moschus ssp.
	Tragulidae	Hyemoschus ssp., Tragulus-Moschiola ssp.
I		

Part II:

- (1) Keep as appropriate.
- (2) This attestation is only applicable to Bovidae and Cervidae.
- (3) This attestation is only applicable to Bovidae and Cervidae other than African buffalo (Syncerus caffer).
- (4) This attestation is only applicable to African buffalo (Syncerus caffer).
- (5) Vaccination is not compulsory, but if the animals have been vaccinated, information on the vaccine(s) used and the time of vaccination shall be
- (6) Date of loading. Imports of these animals shall not be allowed when the animals were loaded either prior to the date of authorisation for exportation to the Union of the third country, territory or part thereof described in Boxes I.7. and I.8., or during a period where restrictive measures have been adopted by the Union against imports of these animals from that country territory or part thereof.

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUN	NTRY			Model RUM-A
II.	Health information	II.a.	Certificate reference number	II.b.
Offici	ial veterinarian			
Name (in capital letters): Qualification and title:				ion and title:
	Date:	Signature:		
8	Stamp:			
	Model S	UI-A		
JNTRY	1			Veterinary certificate to EL
1.1.	Consignor	1.2.	Certificate reference No	1.2.a.
	Name			

COL I.3. Central competent authority Address Tel. I.4. Local competent authority Part I: Details of dispatched consignment Consignee I.6. Name Address Postal code I.7. Country of origin ISO code I.8. Region of origin 1.9. Country of ISO code I.10. Region of destination Code Code destination I.11. Place of origin I.12. Name Address Approval number I.13. Place of loading I.14. Date of departure Approval number I.15. Means of transport I.16. Entry BIP in EU Aeroplane Ship 🔲 Railway wagon Other Road vehicle l.17. Identification Documentary references I.19. Commodity code (HS code) **01.06.19** I.18. Description of commodity I.20. Quantity I.21. I.22. Number of packages I.23. Seal/Container No 1.24. I.25. Commodities certified for: Approved body 1.26. I.27. For import or admission into EU I.28. Identification of the commodities Identification system Identification number Species Age Sex (scientific name)

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model SUI-A

Health information II.a. Certificate reference number II.b.

II.1. Animal health attestation

II.

Part II: Certification

I, the undersigned official veterinarian responsible for the approved body, institute or centre/holding (1) of origin certify that the animals described in Part I meet the following requirements:

- II.1.1. They come from the country, territory or part thereof described in Box I.7.
 - (a) where the diseases referred to in this certificate are notifiable,
 - (b) which at the date of issuing this certificate has been free for the past 12 months from rinderpest.
- II.1.2. They come from the body, institute or centre/holding (1) described in Box I.11.
 - (a) which is approved according to the requirements and conditions set out in Part 3 and 4 of Annex VI to Regulation (EU) No 206/2010;
 - (b) which is not subjected to any restrictions relating to a national programme for the control of infectious diseases to which the animals referred to in Box 1.28, are susceptible;
 - (c) where there have been no clinical cases of the following diseases to which the animals referred to in Box I.28. are susceptible:
 - anthrax for the last 30 days;
 - foot-and-mouth disease, vesicular stomatitis, rabies, African swine fever, classical swine fever and swine vesicular disease for the past 6 months;
 - (d) where there have been no clinical or non-clinical cases of tuberculosis and brucellosis for the past 6 months;
 - (e) around which in an area of radius of 10 km for the last 12 months, there has been no case/outbreak of African swine fever, classical swine fever and swine vesicular disease;
 - (f) around which in an area of 10 km radius for the past 30 days, there has been no case/outbreak of foot-and-mouth disease or vesicular stomatitis.
 - (g) in which they have remained since birth or for the past 6 months before dispatch to the Union.

II.1.3. They:

- (a) have not come into contact with other animals not complying with at least the same health requirements as described in this certificate since birth or for the last 30 days and during their transportation from the approved body, institute or centre/ holding (¹) to the place of shipment;
- (b) were examined by an official veterinarian within 24 hours of loading and showed no clinical sign of disease and are fit for the intended transport;
- (c) are not animals to be killed under a national programme for the eradication of diseases.

II.1.4. Foot-and-Mouth Disease

- either (1) [(a) They come from the country, territory or part thereof described in Box I.7. which at the date of issuing this certificate has been free for the past 12 months from foot-and-mouth disease and;]
- or (¹) [(a) They have been subjected to a virological and serological test for evidence of foot-and-mouth disease virus infection carried out in accordance with one of the prescribed tests for international trade laid down in the OIE Manual of Diagnostic Tests and Vaccines for Terrestrial Animals (OIE Terrestrial Manual), with negative results, taken in the 10 days prior to dispatch to the Union; and]
 - (b) they have not been vaccinated against foot-and-mouth disease.

II.1.5. Brucellosis

- (1) either [They come from the country, territory or part thereof described in Box I.7 which has been free for the past 12 months from brucellosis and have not been vaccinated against that disease]
- (¹)(³) or [They have been subjected, with negative results, to a buffered Brucella antigen test for porcine brucellosis taken in the 30 days prior to dispatch to the Union.]

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model SUI-A Health information II.a. Certificate reference number II.b. II.1.6. Swine vesicular disease (1) either [They come from the country, territory or part thereof described in box 1.7 which has been free for the past 12 months from swine vesicular disease.] (1) or [They have been subjected, with negative results, to a virology and serology test for evidence of swine vesicular disease, as laid down and prescribed for international trade by the OIE Terrestrial Manual, taken in the 30 days prior to dispatch to the Union.] II.1.7. Vesicular Stomatitis (1) either [They come from the country, territory or part thereof described in Box I.7 which has been free for the last 6 months from vesicular stomatitis.] [They have been subjected, with negative results, to a virology and serology test for evidence of vesicular stomatitis, as laid down and prescribed for international trade by the OIE Terrestrial Manual, taken in the 30 days prior to dispatch to the Union.] (1) or II.1.8. Classical swine fever (1) either [They come from the country, territory or part thereof described in Box I.7 which has been free for the past 12 months from (1) or [They have been subjected to a virological and serological test for classical swine fever carried out in accordance with one of the prescribed tests for international trade laid down in the OIE Terrestrial Manual, with negative results, taken in the 30 days prior to dispatch to the Union.] II.1.9. African swine fever (1) either [They come from the country, territory or part thereof described in Box I.7 which has been free for the past 12 months from [They have been subjected, with negative results, to a virus and serology test for African swine fever, as laid down and prescribed for international trade in the OIE Terrestrial Manual, taken in the 30 days prior to dispatch to the Union.] II.1.10. Aujeszky's disease According to official information, no clinical, pathological or serological evidence of Aujeszky's disease has been recorded for the last 12 months in the approved body, institute or centre/holding (1) and in an area with a 5 km radius around the approved body, centre or institute, and They have been subjected, with negative results, to a virology and serology test for evidence of Aujeszky's disease, as laid down and prescribed for international trade by the OIE Terrestrial Manual, taken in the 30 days prior to dispatch to the Union, They have not been vaccinated against Aujeszky's disease and have not been in contact with vaccinated animals. II.1.11. Other vaccinations (a) They have not been vaccinated against rinderpest, vesicular stomatitis, classical swine fever or swine vesicular disease, (2)(b) They have been vaccinated against: II.1.12. Parasite treatment

They have been treated at least twice in the 40 days prior to dispatch to the Union against internal and external parasites with the following product(s) Specify the active ingredients and the doses of the products used

Stamp:

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNT	RY				Model SUI-A
II.	Health info	ormation		II.a. Certificate reference number	II.b.
	II.1.13.	Loading on the mea	ns of transport		
		described in Box I.15	5. that were cleaned and disir	dd/mm nfected before loading with an official on of flow or fall out of the vehicle	ally authorised disinfectant and so
Notes					
				l. 28. coming from an approved body, centre located within a Member State.	
Part I:					
— Вох	reference I			er and lorries), flight number (aircraft) shall inform the BIP of entry into the	
— Вох	reference I			system (tag, tattoos, brand, chip, trans mit tracing of their premises of origin.	ponder). The identifier shall include
		Age: months.			
		Sex (M = male,	F = female, C = castrated).		
		Species Select	the species amongst those liste	ed below:	
Order		Family	Genera/species		
Artioda	ctyla	Suidae	Babyrousa ssp., Hylochoerus	ssp., Phacochoerus ssp., Potamocho	erus ssp., Sus ssp.
		Tayassuidae	Catagonus ssp., Pecari-Tayas	su ssp.	
		Hippopotamidae	Hexaprotodon-Choeropsis, Hip	ppopotamus ssp.	
Part II:					
(1) Kee	p as appro	priate.			
	2) Vaccination is not compulsory, but if the animals have been vaccinated, information on the vaccine(s) used and the time of vaccination must be filled in.				
	³) Tests carried out in accordance with the protocols that, for the disease concerned, are described in Part 6 of Annex I to Regulation (EU) No 206/2010.				
exp	4) Date of loading. Imports of these animals shall not be allowed when the animals were loaded either prior to the date of authorisation for exportation to the Union of the country, territory or part thereof decribed in Boxes I.7. and I.8., or during a period where restrictive measures have been adopted by the Union against imports of these animals from that country,territory or part thereof.				
Official	veterinariar	1			
Nan	ne (in capita	al letters):		Qualifica	ation and title:
Date	е:			Signatur	e:

Status: Point in time view as at 01/07/2016. **Changes to legislation:** There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

Model TRE-A

COUNTRY Veterinary certificate to						ry certificate to EU
	l.1.	Consignor Name	I.2. Certificat	te reference No	1.2.a.	
		Address	I.3. Central competent authority			
	Tel.			mpetent authority	<i>y</i>	
nent			I.4. Local co	mpotont dutions	,	
sign	1.5.	Consignee Name	1.6.			
con						
hed		Address Postal code				
patc		Tel.				
fdis	1.7.	Country of origin ISO code I.8. Region of origin Code	I.9. Country			
ils o			destination	on 	destination	on
Partl: Details of dispatched consignment	l.11.	Place of origin	I.12.	I		
₩.		Name Approval number				
Ра		Address				
	I.13.	Place of loading Address Approval number	I.14. Date of	departure		
	l.15.	Means of transport	I.16. Entry Bli	P in EU		
	Aeroplane Ship Railway wagon Road vehicle Other					
		Identification	l.17.			
		Documentary references				
	I.18.	Description of commodity		I.19. Commodit	ty code (HS code) 01.06.19	
					I.20. Quantity	
	I.21.				I.22. Number of p	ackages
	1.23.	Seal/Container No			1.24.	
	1.25.	Commodities certified for:				
		Approved body				
	1.26.		I.27. For impo	ort or admission i	into EU	
	1.28.	Identification of the commodities	l			
		Species Identification system (scientific name)	Identification	number	Age	Sex

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY Model TRE-A

II. Health information II.a. Certificate reference number II.b.

II.1. Animal health attestation

Part II: Certification

I, the undersigned official veterinarian responsible for the approved body, institute or centre/holding (1) of origin certify that the animals described in Part I meet the following requirements:

- II.1.1. They come from the third country, territory or part thereof described in Box I.7.
 - (a) where the diseases referred to in this certificate are notifiable,
 - (b) which at the date of issuing this certificate has been free for the past 12 months from rinderpest.
- II.1.2. They come from the body, institute or centre/holding (1) described in Box I.11.,
 - (a) which is approved according to the requirements and conditions set out in Part 3 and 4 of Annex VI to Regulation (EU) No 206/2010;
 - (b) which is not subjected to any restrictions relating to a national programme for the control of infectious diseases to which the animals referred to in Box 1.28, are susceptible;
 - (c) where there have been no clinical cases of the following diseases to which the animals referred to in Box I.28. are susceptible:
 - anthrax for the last 30 days;
 - foot-and-mouth disease, rabies, (1)(2) [African horse sickness] for the past 6 months,
 - (d) where there have been no clinical or non-clinical cases of tuberculosis for the past 6 months;
 - (e) around which in an area of 10 km radius for the last 30 days, there has been no case/outbreak of foot-and-mouth disease,
 - (f) in which they have remained since birth or for the past 6 months before dispatch to the Union,
- (1)(2) [(g) around which in an area of radius of 150 km for the last 60 days, there has been no case/outbreak of African horse sickness].
- II.1.3. They:
 - (a) have not come into contact with other animals not complying with at least the same health requirements as described in this
 certificate since birth or for the past 30 days and during their transportation from the approved body, institute or centre/holding (¹) to the place of shipment;
 - (b) were examined by an official veterinarian within 24 hours of loading and showed no clinical sign of disease and are fit for the intended transport:
 - (c) are not animals to be killed under a national programme for the eradication of diseases.

(1)(3) [II.1.4. Foot-and-Mouth Disease

- either (1) [(a) They come from the country, territory or part thereof described in Box I.7 which has been free for the past 12 months from foot-and-mouth disease with or without vaccination, and]
- or (1) [(a) They have been subjected to the following tests:
 - a serological test for evidence of foot-and-mouth disease virus infection carried out in accordance with one of the
 prescribed tests for international trade laid down in the OIE Manual of Diagnostic Tests and Vaccines for Terrestrial
 Animals (OIE Terrestrial Manual), with negative results, taken in the 10 days prior to dispatch to the Union, and
 - [a probang test for evidence of foot-and-mouth disease virus infection carried out in accordance with the procedures
 described in the OIE Terrestrial Manual with negative results, taken 10 days prior to dispatch to the Union and]
 - (b) have not been vaccinated against foot-and-mouth disease.

II.1.5. Other vaccinations

(a) They have not been vaccinated against rinderpest,

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNT	RY					Model TRE-
II.	Health in	formati	ion		II.a. Certificate reference number	II.b.
	(4) (b) They have been vaccinated against:					
			[anthrax on the .used)],	(dd/mm/yyyy)(d	ate(s)) with the following vaccine(s)	(name of vaccine(s
		(¹) [rabies on the	(dd/mm/yyyy)(date(s))) with the following vaccine(s)	(name of vaccine (s) used)]
	II.1.6.	Para	site treatment			
					prior to dispatch to the Union against active ingredients and the doses of t	
	II.1.7.	Load	ding on the mea	ans of transport		
		desc	ribed in Box I.1	5 that were cleaned and disir	n(dd/mm nfected before loading with an officia d not flow or fall out of the vehicle	ally authorised disinfectant and so
Notes						
					28. coming from an approved body, ir centre located within a Member Stat	
Part I:						
— Вох	reference	l.15.:			ner and lorries), flight number (aircraft) r shall inform the BIP of entry into the	
— Вох	reference	l.28.:			system (tag, tattoos, brand, chip, trans mit tracing of their premises of origin.	ponder). The identifier shall include
			Age: months.			
			Sex (M = male	, F = female, C = castrated).		
			Species: Select	t the species amongst those list	ted below:	
Order		Fa	mily	Genera/species		
Perisso	dactyla	Ta	piridae	Tapirus ssp.		
		Rh	inocerotidae	Ceratotherium ssp., Dicerorhir	nus ssp., Diceros ssp., Rhinoceros ssp	
Probos	cidea	Ele	phantidae	Elephas ssp., Loxodonta ssp.		
Part II:						
(1) Kee	p as app	opriate				
(²) This	attestation	on is or	nly applicable to	Rhinocerotidae.		
(3) This	attestation	on is or	nly applicable to	Elephas. ssp.		
	cination is d in.	not co	empulsory, but if t	he animals have been vaccinate	ed, information on the vaccine(s) used	and the time of vaccination must be
exp	ortation to	the U	Inion of the third	I country,territory or part thereo	en the animals were loaded either proof described in Boxes I.7. and I.8., o	r during a period where restrictive

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

COUNTRY	Model TRE-A		
II. Health information	II.a. Certificate reference number	II.b.	
Official veterinarian			
Name (in capital letters):	Qualification and title:		
Date:	Signature:		
Stamp:			

PART 3

Requirements concerning bodies, institutes or centres in third countries

The body, institute or centre in a third country must:

- (a) be clearly demarcated and separated from its surroundings;
- (b) have adequate means for catching, confining and isolating animals, and have available adequate quarantine facilities and approved standard operating procedures for animals coming from unknown origin;
- (c) have a vector-protected structure complying with the following requirements:
 - (i) it has appropriate physical barriers at entry and exit points;
 - (ii) the openings of the vector-protected structure are vector-screened with mesh of appropriate gauge impregnated regularly with an approved insecticide according to the instructions of the manufacturer;
 - (iii) vector surveillance and control are carried out within and around the vector-protected structure;
 - (iv) measures are taken to limit or eliminate breeding sites for vectors in the vicinity of the vector-protected structure;
 - (v) standard operating procedures are in place, including descriptions of backup and alarm systems, for the operation of the vector-protected structure and for the transport of the animals from that structure to the place of loading;
- (d) keep, for a minimum period of ten years, up-to-date records indicating:
 - (i) the number and identity (age, sex, species and individual identification, where appropriate) of the animals of each species present on their premises;
 - (ii) the number and identity (age, sex, species and individual identification where appropriate) of animals arriving in or leaving their premises, together with information on their origin or destination, the means of transport, and the health status of those animals;
 - (iii) the results of blood tests or any other diagnostic procedures carried out on the animals on their premises;
 - (iv) cases of disease and, where appropriate, the treatment administered;

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

- (v) the results of the post-mortem examinations on animals that have died on their premises, including still-born animals;
- (vi) observations made during any isolation or quarantine period;
- (e) be free from the diseases listed in Annex A to Directive 92/65/EEC or mentioned in the veterinary certificates for the relevant species set out in Part 2 of Annex VI to this Regulation, for at least the previous three years, as evidenced by the records kept pursuant to point (d) and the results of the clinical and laboratory tests carried out on the animals on their premises;
- (f) either have an arrangement with a laboratory approved by the competent authority to perform post-mortem examinations, or have one or more appropriate premises where these examinations may be performed under the authority of the approved veterinarian;
- (g) ensure disposal of the carcasses of animals which die of a disease or are euthanised;
- (h) secure, by contract or legal instrument, the services of a veterinarian approved by and acting under the control of the competent authority, who must perform at least the following tasks:
 - (i) ensure that appropriate disease surveillance and control measures are applied in that body, institute or centre. Such measures must be approved by the competent authority of the third country, territory or part thereof where the body, institute or centre is situated, taking into account the disease situation and must include at least the following elements:
 - an annual disease surveillance plan including appropriate control measures concerning zoonoses in the animals present on the premises,
 - clinical, laboratory and post-mortem testing of animals suspected to be affected by transmissible diseases and zoonoses,
 - vaccination of susceptible animals against infectious diseases and zoonoses;
 - (ii) ensure that any suspect deaths or the presence of any other symptom suggesting that animals have contracted one or more of the diseases listed in Annex A to Directive 92/65/EEC or mentioned in the veterinary certificates for the relevant species set out in Part 2 of Annex VI to this Regulation are notified without delay to the competent authority, where that particular disease is notifiable in the third country, territory or part thereof concerned;
 - (iii) ensure that incoming animals have been quarantined as necessary, in accordance with the instructions given by the competent authority;
 - (iv) ensure compliance with the animal health requirements which the animals must fulfil in order to be introduced into the Union.

PART 4

Conditions concerning the approval of bodies, institutes or centres in third countries

1. Approval must be granted only to those bodies, institutes or centres which comply with the requirements set out in Part 3.

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

- 2. Where vector protection is required, the approval of a structure as vector-protected must be granted only if the criteria in point (c) of Part 3 are met. In order to grant the approval, the competent authority must verify at least three times during the required protection period (at the beginning, during and at the end of the period) the effectiveness of the vector protection measures, by means of a vector trap inside the vector protected structure.
- 3. Each approved body, institute and centre must be assigned an approval number.
- 4. Approval must be maintained only as long as the following conditions continue to be met:

the premises are under the control of an official veterinarian, who must perform at least the following tasks:

- (i) inspect the premises of the body, institute or centre at least once per year;
- (ii) audit the activity of the veterinarian referred to in point (h) of Part 3 and the implementation of the annual disease surveillance plan referred to in the first indent of point (h)(i);
- (iii) ensure that the provisions laid down in Parts 3 and 4 are met;
- (iv) verify that:
 - compliance with the animal health requirements which the animals must fulfil in order to be introduced into the Union;
 - the results of the clinical, post-mortem and laboratory tests on the animals have revealed no occurrence of the diseases listed in Annex A to Directive 92/65/EEC or mentioned in the veterinary certificates for the relevant species set out in Part 2 of Annex VI to this Regulation.
- 5. The approval must be withdrawn where the competent authority finds that the requirements of Part 3 are no longer being fulfilled.
- 6. Where notification is given of the suspicion of the occurrence of one of the diseases listed in Annex A to Directive 92/65/EEC or mentioned in the veterinary certificates for the relevant species laid down in Part 2 of Annex VI to this Regulation, the competent authority must suspend the approval of the body, institute or centre, until the suspicion has been officially ruled out. Depending on the disease involved and the risk of disease transmission, the suspension may relate to the the body, institute or centre as a whole or only to certain categories of animals susceptible to the disease in question. The competent authority must ensure that the measures necessary to confirm or rule out the suspicion and to avoid any spread of disease are taken.
- 7. Where the suspected disease referred to in point 6 is confirmed, the approval of the body, institute or centre must be withdrawn.
- 8. Where the approval of a body, institute or centre has been withdrawn, it must be restored only where the following conditions are complied with:
- (a) the disease and the source of infection were eradicated on the premises of the body, institute or centre concerned;
- (b) the premises of the body, institute or centre concerned were appropriately cleaned and desinfected;

Status: Point in time view as at 01/07/2016.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

- (c) the body, institute or centre concerned complies with the requirements set out in points (a) to (d) and (f) to (h) of Part 3.
- 9. The competent authority which approved the body, institute or centre must inform the Member States that included the body, institute or centre on their lists of approved bodies, institutes and centres of the suspension, withdrawal or restoration of that approval.]

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010. (See end of Document for details)

- (1) $[^{X1}[^{F1}OJ 121, 29.7.1964, p. 1977/64.]]$
- (2) $[^{X1}[^{F1}OJ L 46, 19.2.1991, p. 19.]]$
- (3) [X1]F1Delete country as applicable.]]
- (4) [X1[F1Serbia, not including Kosovo under UNSCR 1244/99.]]
- (5) [X1OJ L 249, 23.7.2004, p. 20.]
- **(6)** [X1OJ L 59, 4.3.2008, p. 19.]
- (7) [X1OJ L 167, 7.7.2000, p. 22.]
- (8) [X1OJ L 39, 9.2.2002, p. 71.]
- (9) [X1OJ L 268, 24.9.1991, p. 56.]
- (10) [X1OJ L 13, 16.1.1997, p. 28.]

Editorial Information

X1 Substituted by Corrigendum to Commission Regulation (EU) No 206/2010 of 12 March 2010 laying down lists of third countries, territories or parts thereof authorised for the introduction into the European Union of certain animals and fresh meat and the veterinary certification requirements (Official Journal of the European Union L 73 of 20 March 2010).

Textual Amendments

F1 Substituted by Commission Implementing Regulation (EU) No 644/2012 of 16 July 2012 amending Regulation (EU) No 206/2010 laying down lists of third countries, territories or parts thereof authorised for the introduction into the European Union of certain animals and fresh meat and the veterinary certification requirements, as regards Russia (Text with EEA relevance).

Status:

Point in time view as at 01/07/2016.

Changes to legislation:

There are currently no known outstanding effects for the Commission Regulation (EU) No 206/2010.