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▶<u>B</u> 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)

(OJ L 219, 15.8.2013, p. 1)

Corrected by:

▶<u>C1</u> Corrigendum, OJ L 238, 6.9.2013, p. 23 (780/2013)

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)

THE EUROPEAN COMMISSION,

Having regard to the Treaty on the Functioning of the European Union,

Having regard to Council Directive 2004/68/EC of 26 April 2004 laying down animal health rules for the importation into and transit through the Union of certain live ungulate animals, amending Directives 90/426/EEC and 92/65/EEC and repealing Directive 72/462/EEC (¹), and in particular the first and second subparagraphs of Article 3(1), the first subparagraph of Article 6(1), Article 7(e), Article 8(c) and Article 13(1) thereof,

Whereas:

- (1) Commission Regulation (EU) No 206/2010 (²) lays down the requirements for the introduction into the Union of, amongst others, certain ungulates. That Regulation does not apply to non-domesticated animals intended for an approved body, institute or centre as defined in Article 2(1)(c) of Directive 92/65/EEC of 13 July 1992 laying down animal health requirements governing trade in and imports into the Community of animals, semen, ova and embryos not subject to animal health requirements laid down in specific Community rules referred to in Annex A (I) to Directive 90/425/EEC (³).
- (2) The lack of specific animal health requirements for the introduction of ungulates intended for an approved body, institute or centre into the Union causes practical problems for such structures and strongly limits their activities, because of their need to introduce those animals.
- (3) It is appropriate to lay down animal health requirements for the introduction of ungulates intended for an approved body, institute or centre into the Union, which would take account of the specific situation of such animals. In the interest of simplification of Union law, it is appropriate that such rules be laid down in Regulation (EU) No 206/2010. The scope of that Regulation should therefore be amended accordingly.
- (4) Regulation (EU) No 206/2010 provides that consignments of ungulates may only be introduced into the Union if they come from the third countries, territories or parts thereof listed in Part 1 of Annex I to that Regulation.

^{(&}lt;sup>1</sup>) OJ L 139, 30.4.2004, p. 321.

⁽²⁾ OJ L 73, 20.3.2010. p. 1.

^{(&}lt;sup>3</sup>) OJ L 268, 14.9.1992, p. 54.

- (5) Council Directive 2002/99/EC of 16 December 2002 laying down the animal health rules governing the production, processing, distribution and introduction of products of animal origin for human consumption (¹) provides for the adoption by the Commission of lists of the third countries or regions of third countries from which imports of specified products of animal origin are permitted.
- (6) Council Directive 2009/156/EC of 30 November 2009 on animal health conditions governing the movement and importation from third countries of equidae (²) provides that the importation of equidae into the Union is to be authorised only from third countries that appear on a list to be drawn up or amended in accordance with the procedure laid down in that Directive.
- (7) Council Directive 2009/158/EC of 30 November 2009 on animal health conditions governing intra-Community trade in, and imports from third countries of, poultry and hatching eggs (³) provides that poultry and hatching eggs imported into the Union must have originated in a third country or part of a third country included on a list drawn up by the Commission in accordance with the procedure laid down in that Directive.
- (8) The introduction of ungulates intended for an approved body, institute or centre into the Union should, in particular, fulfil the general requirements for the introduction of live animals into the Union and further specific animal health requirements and offer specific guarantees ensuring that the animals introduced into the Union do not endanger the animal health status of the Union.
- (9) The general requirements for the introduction of live animals into the Union, consisting of an effective system of veterinary services in charge of the control of animal health, are currently fulfilled by those third countries, territories and parts thereof listed pursuant to Directives 2002/99/EC, 2009/156/EC and 2009/158/EC.
- (10) However, the general requirements for the introduction of live animals into the Union do not guarantee that ungulates are free of diseases. Individual animals may still carry infectious diseases that could spread into the Union and consequently consitute a danger to animal health in the Union. Ungulates intended for an approved body, institute or centre should therefore only be introduced into the Union directly from a body, institute or centre which complies with certain requirements and is approved by the competent authority of the third country, territory or part thereof in which it is situated.
- (11) It is appropriate that the list of such bodies, institutes or centres be established by the Member State of destination, following an assessment of all relevant information.

⁽¹⁾ OJ L 18, 23.1.2003, p. 11.

^{(&}lt;sup>2</sup>) OJ L 192, 23.7.2010, p. 1.

⁽³⁾ OJ L 343, 22.12.2009, p. 74.

- (12) In order to protect the animal health in the Union, it is crucial that consignments of ungulates introduced into the Union and destined to approved bodies, institutes or centres be transported directly and without delay to their destination in sealed containers and that further movement of such animals within the Union be restricted.
- (13) In order to address exceptional circumstances such as situations concerning animal welfare problems, conservation of endangered species, sudden natural disasters or political unrest, in which it is not possible to apply all the animal health requirements and especially those related with the approval of the body, institute or centre of origin, Member States should be able to introduce into their territory certain ungulates destined to an approved body, institute or centre, under specific conditions. However, even in such cases, a permit should be required in order to ensure sufficient reduction of the animal health risk.
- (14) Regulation (EU) No 206/2010 should therefore be amended accordingly.
- (15) The measures provided for in this Regulation are in accordance with the opinion of the Standing Committee on the Food Chain and Animal Health,

HAS ADOPTED THIS REGULATION:

Article 1

Regulation (EU) No 206/2010 is amended as follows:

- (1) In Article 1, paragraph 3 is deleted.
- (2) The following Article 3a is inserted:

'Article 3a

Conditions for the introduction of ungulates intended for an approved body, institute or centre

1. By way of derogation from Article 3, the competent authority of a Member State may authorise the introduction into its territory of consignments of ungulates of the species listed in Tables 1, 2 and 3 of Part 1 of Annex VI where those consignments are destined for an approved body, institute or centre, provided that the following conditions are complied with:

- (a) an assessment has been carried out by the competent authority of the Member State of destination of the animal health risks that each of the consignments may present for the Union;
- (b) the consignments concerned come from a third country, territory or part thereof which is included in one of the lists set out in:
 - (i) Part 1 of Annex I or in Part 1 of Annex II to this Regulation,

- (ii) Decision 2004/211/EC (*), Decision 2007/777/EC (**), Regulation (EC) No 798/2008 (***), Regulation (EC) No 119/2009 (****), Regulation (EU) No 605/2010 (*****),
- (c) the ungulates originate from a body, institute or centre in a third country, territory or part thereof, referred to in point (a), which is included in a list established in accordance with Article 3c;
- (d) the ungulates have been quarantined in a vector-protected facility at the premises of the body, institute or centre referred to in point (c) for the period provided for in the relevant certificates;
- (e) the ungulates are conveyed directly to an approved body, institute or centre in the Member State of destination;
- (f) the ungulates are accompanied by an appropriate veterinary certificate, drawn up in accordance with the relevant model of veterinary certificate referred to in Tables 1, 2 and 3 in Part 1 of Annex VI and set out in Part 2 of that Annex;
- (g) the ungulates comply with the requirements set out in the model of veterinary certificate referred to in point (f).

The Member State of destination shall inform the Commission and the other Member States in the Standing Committee on the Food Chain and Animal Health of the authorisation granted pursuant to the first subparagraph, prior to the introduction of the ungulates into their territory.

2. Where exceptional circumstances render compliance with points (c) and (d) of paragraph 1 impossible, the competent authority of the Member State of destination may authorise the introduction, into its territory, of ungulates of the species listed in Tables 1, 2 and 3 of Part 1 of Annex VI from *other holdings* which do not comply with the requirements laid down in those points, provided that the requirements laid down in points (a), (b) and (e) to (g) of paragraph 1 are complied with and that the following additional conditions are met:

- (a) a prior application for a permit has been made by the owner, or a natural person representing that owner, and the Member State of destination has granted such permit after having carried out a risk assessment that has indicated that the introduction of the ungulates concerned into its territory does not constitute an animal health risk for the Union;
- (b) the ungulates have been quarantined in the third country, territory or part thereof of origin under official supervision for the time necessary for them to meet the animal health conditions set out in the model of veterinary certificate referred to in point (f):
 - (i) at a place approved by the competent authority of the third country, territory or part thereof of origin of the animals;
 - (ii) in accordance with the arrangements prescribed in the permit that shall provide at least the same guarantees as those laid down in points (a), (b) and (e) to (g) of paragraph 1.

Where ungulates are introduced into the Union pursuant to the first subparagraph, they shall be quarantined in an approved body, institute or centre *of destination* for at least six months from the time of introduction into the Union, during which period the requirements provided for in Article 8(1)(a) of Council Directive 90/425/EEC may be applied by the competent authorities.

The Member State authorising the introduction of ungulates pursuant to the first subparagraph shall inform the Commission and the other Member States in the Standing Committee on the Food Chain and Animal Health of such authorisation, prior to the introduction of the ungulates into its territory.

(*) OJ L 73, 11.3.2004, p. 1. (**) OJ L 312, 30.11.2007, p. 49. (***) OJ L 226, 23.8.2008, p. 1. (****) OJ L 39, 10.2.2009, p. 12. (*****) OJ L 175, 10.7.2010, p. 1.'

(3) The following Article 3b is inserted:

'Article 3b

Conditions for the entry and transit of ungulates intended for an approved body, institute or centre through the territory of Member States other than the Member State of destination

The transit of the ungulates referred to in Article 3a through a Member State other than the Member State of destination shall be permitted only subject to the authorisation of the competent authority of the Member State of transit. Such authorisation may be granted only on the basis of a risk assessment by that competent authority, in view of the information submitted to it by the Member State of destination.

The Member State of destination shall inform the Commission and the other Member States in the Standing Committee on the Food Chain and Animal Health, prior to the transit, when authorising the introduction of animals under the conditions provided for in Article 3a.'

(4) The following Article 3c is inserted:

'Article 3c

List of approved bodies, institutes or centres in third countries, territories and parts thereof

1. Following an assessment of compliance with the conditions laid down in paragraph 2, each Member State may establish a list of bodies, institutes and centres from which the introduction of ungulates into its territory may be authorised pursuant to Article 3a(1).

2. A body, institute or centre in a third country, territory or part thereof shall only be included in the list referred to in paragraph 1 where the following conditions are complied with:

- (a) the body, institute or centre complies with the requirements set out in Part 3 of Annex VI;
- (b) the body, institute or centre is approved by the competent authority of the third country, territory or part thereof where that body, institute or centre is situated;

(c) the competent authority of the third country, territory or part thereof provides sufficient guarantees that the conditions concerning the approval of bodies, institutes or centres set out in Part 4 of Annex VI are complied with.

3. A Member State may include in the list referred to in paragraph (1) bodies, institutes or centres in third countries which are already included in such a list established by another Member State, without having assessed compliance with the conditions laid down in paragraph 2.

4. Member States shall keep the lists referred to in paragraph (1) up to date, taking into account in particular any suspension or withdrawal of the approval granted by the competent authority of a third country, territory or part thereof to the bodies, institutes or centres situated therein and included in those lists.

5. Member States shall make available to the public, by means of Internet-based information pages, the lists referred to in paragraph 1 and shall keep those Internet-based information pages up to date.

6. Member States shall communicate the Internet address of their Internet-based information pages to the Commission.'

(5) Article 4 is replaced by the following:

'Article 4

Conditions for the assembly centres for certain consignments of ungulates

1. Consignments of ungulates which contain live animals from more than one holding shall only be introduced into the Union if they are assembled in assembly centres approved by the competent authority of the third country, territory or part thereof of origin of the animals in accordance with the requirements set out in Part 5 of Annex I.

2. Consignments of ungulates introduced into the Union in accordance with Article 3a or Article 6 shall not originate from more than one holding and shall not be assembled in assembly centres.'

- (6) In Article 8, point (b) is replaced by the following:
 - '(b) unloaded in, or when transported by air, moved to another aircraft, or transported by road, by rail, or moved on foot through a third country, territory or part thereof which is not authorised for imports of the animals concerned into the Union.'

(7) In Article 11, paragraph 1 is replaced by the following:

'1. Following their introduction into the Union, consignments of ungulates, other than those referred to in Article 3a shall be conveyed in a vector-protected means of transport without delay to the holding of destination.

Those ungulates shall remain on that holding for a period of at least 30 days, unless they are dispatched directly to a slaughterhouse.'

(8) The following Article 13a is inserted:

'Article 13a

Conditions to be applied following the introduction of consignments of ungulates intended for approved bodies, institutes or centres

1. Following their introduction into the Union, consignments of ungulates intended for approved bodies, institutes or centres shall be transported without delay to the approved body, institute or centre of destination in means of transport that are vector-protected and so constructed that the animals cannot escape and faeces, urine, litter, fodder, waste or any other material cannot flow or fall out from the vehicle or container during transportation.

2. The animals shall be kept in quarantine in vector-protected facilities on the premises of the approved body, institute or centre of the Member State of destination for a minimum of 30 days. After the 30 days quarantine period the animals may be moved to another approved body, institute or centre.

3. Animals introduced into an approved body, institute or centre can only be moved to a destination other than an approved body, institute or centre provided that:

- (a) at least six months have elapsed from the time of introduction into the Union, and
- (b) the movement is carried out in accordance with paragraph 4 of Annex C to Directive 92/65/EEC.

4. By way of derogation from paragraph 3, animals may leave an approved body, institute or centre before the end of the six-month period provided for in that paragraph, only where the following conditions are complied with:

- (a) the animals are exported to a third country, territory or part thereof;
- (b) for the purpose of their export as referred to in a) the animals are transported in means of transport that are vector-protected and so constructed that the animals cannot escape and faeces, urine, litter, fodder, waste or any other material cannot flow or fall out from the vehicle or container during transportation.'
- (9) Annex VI, the text of which is set out in the Annex to this Regulation, is added.

Article 2

Entry into force

This Regulation shall enter into force on the twentieth day following that of its publication in the *Official Journal of the European Union*.

This Regulation shall be binding in its entirety and directly applicable in all Member States.

ANNEX

'ANNEX VI

PART 1

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Table 1				
		rtificate for animals of the species listed below that are tended for an approved body, institute or centre.		
Order				
Artiodactyla	Antilocapridae	Antilocapra ssp.		
	Bovidae	Addax ssp., Aepyceros ssp., Alcelaphus ssp., Ammo- dorcas ssp., Ammotragus ssp., Antidorcas ssp., Antilope ssp., Bison ssp., Bos ssp. (including Bibos, Novibos, Poephagus), Boselaphus ssp., Bubalus ssp. (including anoa), Budorcas ssp., Capra ssp., Cepha- lophus ssp., Connochaetes ssp., Damaliscus ssp. (including Beatragus), Dorcatragus ssp., Gazella ssp., Hemitragus ssp., Madoqua ssp., Naemorhedus ssp., Litocranius ssp., Madoqua ssp., Naemorhedus ssp. (including Nemorhaedus and Capricornis), Neotragus ssp., Oreamnos ssp., Oreotragus ssp., Oryx ssp., Ourebia ssp., Pocapra ssp., Pseudois ssp., Pseudoryx ssp., Raphicerus ssp., Redunca ssp., Rupicapra ssp., Saiga ssp., Sigmoceros-Alecelaphus ssp., Sylvicapra ssp., Tragelaphus ssp. (including Booce- rus).		
	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						
	Model of veterinary certificate for animals of the species listed below that are originating from and intended for an approved body, institute or centre.					
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': Model of veterinary certificate for animals of the species listed below that are originating from and intended for an approved body, institute or centre.					
Order	Family	Genera/species			
Perissodactyla	Tapiridae	Tapirus ssp.			
	Rhinocerotidae	Ceratotherium ssp., Dicerorhinus ssp., Diceros ssp., Rhinoceros ssp.			
Proboscidea	Elephantidae	Elephas ssp., Loxodonta ssp.			

PART 2

			Model RU	M-A					
ου	NTR					•		erinary ce	ertificate to E
	1.1.	Consignor Name		1.2. Cer	tificate refer	ence No	1.2.a.		
				I.3. Cer	ntral compet	tent authority			
		Address Tel.				,			
₌│		Tel.		I.4. Loc	al compete	nt authority			
₽	1.5	0		1.0					
<u>_</u>	1.5.	Consignee Name		1.6.				_	
Į									
		Address Restal cada							
5		Postal code Tel.							
	1.7.	Country of origin ISO code I	.8. Region of origin Code		untry of tination	ISO code		on of nation	Code
2									
ġ	1.11.	Place of origin		1.12.					
ן <u>נ</u>		<u> </u>					_		
		Name	Approval number						
-		Address							
╞	1.40								
	1.13.	Place of loading Address	Approval number	1.14. Dat	e of departi	ure			
		Addicas							
1	l.15.	I.15. Means of transport		I.16. Entry BIP in EU					
		Aeroplane Ship Road vehicle Other	Railway wagon 🔲						
		Identification		1.17.					
		Documentary references							
ł	l.18.	Description of commodity			l.19.	Commodity	code (HS cod	e)	
		,						-,	
						1	I.20. Quantity		
╞	1.01								
	1.21.						I.22. Number	от расказ	jes
	1.23.	Seal/Container No					1.24.		
	1.25.	Commodities certified for:							
		Approved body							
ł	1.26.			1.27. For	import or a	dmission int	o EU		
								_	
	1.28.	Identification of the commodities							
		Species	Identification exetem	- اما	ntification	Imbor	A		Ser
		Species (scientific name)	Identification system	Ide	ntification n	INDE	Age		Sex
- L									

	COUNT	RY		Model RUM-A					
	II.	Health info	ormation	II.a. Certificate reference number	II.b.				
	II.1.	Animal he	ealth attestation						
		I, the und described	dersigned official veterinarian responsible for the approved body, institute or centre/holding (1) of origin certify that the animals d 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 a (b) which at the date of issuing this certificate has be						
ation	st. <								
ertific		II.1.2.	They come from the body, institute or centre/holding (⁽¹⁾ described in Box I.11;					
Part II: Certification			 (a) which is approved according to the requirements No 206/2010; 	and conditions set out in Part 3 and	I 4 of Annex VI to Regulation (EU)				
P			(b) which is not subjected to any restrictions relating to animals referred to in Box I.28. are susceptible;	o a national programme for the contro	l of infectious diseases to which the				
			(c) where there have been no clinical cases of the susceptible:	following diseases to which the ar	imals referred to in Box I.28. are				
			— anthrax for the last 30 days;						
_			 foot-and-mouth disease, bluetongue, Rift valley lumpy skin disease, peste des petits ruminants, 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 las animals referred to in Box I.28. are susceptible: for monia, peste des petits ruminants, sheep pox, go	ot-and-mouth disease, vesicular stoma	titis, contagious bovine pleuropneu-				
			 (f) around which in an area of 150 km radius for the la animals referred to in Box I.28. are susceptible: b disease; 						
			(g) in which they have remained since birth or for the	e past 6 months before dispatch to th	ne Union.				
		II.1.3.	They:						
			 (a) have not come into contact with other animals not certificate for the last 30 days and during their tra place of shipment; 						
			 (b) were examined by an official veterinarian within 24 intended transport; 	hours of loading and showed no clinic	al sign of disease and are fit for the				
			(c) are not animals to be killed under a national prog	ramme for the eradication of disease	s.				
		II.1.4.	Foot-and-Mouth Disease						
		either (1)	[(a) They come from the country, territory or part there foot-and-mouth disease with or without vaccination		en free for the past 12 months from				
		or (1)	[(a) They have been subjected to the following tests:						
			 a serological test for evidence of foot-and-mu prescribed tests for international trade laid do Animals (OIE Terrestrial Manual), with negative 	wn in the OIE Manual of Diagnostic	Tests and Vaccines for Terrestrial				
			 (1)(2)[a probang test for evidence of foot-and-modescribed in the OIE Terrestrial Manual v Union] (1)(4)[taken on two occasions 15 days dispatch to the Union, and] 	vith negative results, (1)(3)[taken 10) days prior to dispatch to the				
	▶	²⁾ (¹)	(b) they have not been vaccinated against foot-and-m	nouth disease.◀					

Health inf	ormation II.a. Certificate reference number II.b.
II.1.5.	Bluetongue and Epizootic haemorrhagic disease (EHD)
either (1)	[They come from the country, territory or part thereof described in Box I.7 which has been free for 24 months from bli tongue/EHD in accordance with the OIE Terrestrial Animal Health Code (OIE Terrestrial Code).]
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 shipment and were subjected to a serology test according to the OIE Terrestrial Manual, with negative results, carried out 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 shipment and were subjected to a PCR test according to the OIE Terrestrial Manual, with negative results, carried out at le 14 days after introduction into the approved body, institute or centre.]
or (1)	[They come from a seasonally free area and were subjected during that period to an serology test according to the C Terrestrial Manual, with negative results, carried out at least 28 days after introduction into the approved body, institute centre/holding ⁽¹⁾ .]
or (1)	[They come from a seasonally free area and were subjected during that period to a PCR test according to the OIE Terrest Manual, with negative results, carried out at least 14 days after introduction into the approved body, institute or centre/hc ing (¹).]
II.1.6.	Rift valley fever
either (¹)	[They come from the country, territory or part thereof described in Box I.7. which has been free for 48 months from Rift val fever and have not been vaccinated against that disease.]
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 shipment during which the animals showed no clinical signs of Rift valley fever and were protected from vectors between i 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 (*) with negative results for evidence of Rift valley fever, as laid do and prescribed for international trade by the OIE Terrestrial Manual, taken at the beginning of the isolation/quarantine period a at least 42 days later on, the second of which must have been taken within 10 days of dispatch to the Union.]
II.1.7.	Brucellosis
either (1)	[They come from a country, territory or part thereof described in Box I.7 which has been free for the past 12 months free brucellosis and which have not been vaccinated against that disease;]
or (1)	[They have been subjected to a test as laid down and prescribed for international trade by the OIE Terrestrial Manual, in the days prior to dispatch to the Union;]
or (1)	[They are castrated males of any age].
II.1.8.	Other vaccinations
	(a) They have not been vaccinated against vesicular stomatitis,
(5)) (b) They have been vaccinated against:
	(¹) [anthrax on the (dd/mm/yyyy)(date(s)) with the following vaccine(s) (name of vaccine used)],
	(¹) [rables on the
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 parasi with the following product(s)
II.1.10.	Loading on the means of transport
	They have been loaded for dispatch to the Union on(dd/mm/yyyy) (⁶) in the means of transport described Box 1.15. that were cleaned and disinfected before loading with an officially authorised disinfectant and so constructed ti faeces, urine, litter or fodder could not flow or fall out of the vehicle or container during transportation.

COUNTI					Model RUM-A		
П.	Health inform	nation		II.a. Certificate reference number	II.b.		
Notes							
					institute or centre in a third country, te. 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, case of unloading and reloading, the consignor shall inform the BIP of entry into the EU.							
— Box	reference 1.19	9.: Use approp	iate HS code: 010613 or 010619.				
- Box reference I.28.: Identification				ystem (tag, tattoos, brand, chip, tran mit tracing of their premises of origi	sponder). The identifier shall include n.		
Age: months			5.				
		Sex (M = m	ale, F = female, C = castrated).				
Species: Select the species amongst those listed below:							
Order		Family	Genera/species				
Artioda	ctyla	Antilocapridae	Antilocapra				
Bovidae		Bovidae	Addax ssp., Aepyceros ssp., Alcelaphus ssp., Ammodorcas ssp., Ammotragus ssp., Antidorcas ssp., Antiope ssp., Bison ssp., Bos ssp. (including Bibos, Novibos, Poephagus), Boselaphus ssp., Bubaliscus 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 Capricomis), Neotragus ssp., Oreatmos ssp., Oreotragus ssp., Ory, ssp., Ourebia ssp., Ovibos ssp., Oris ssp., Patholops ssp., Pelea ssp., Rocapra ssp., Pseudolo; ssp., Pseudoryx ssp., Raphicerus ssp., Tedurca ssp., Rupicapra ssp., Saiga ssp., Sigmoceros-Alecelaphus ssp., Sylvicapra ssp., Syncerus ssp., Taur- otragus ssp., Tetracerus ssp., Tragelaphus ssp. (including Boccerus).				
		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-Mo	schiola ssp.			
Part II:							
(¹) Kee	p as appropri	ate.					
(²) This	attestation is	only applicable t	<i>Bovidae</i> and <i>Cervidae</i> .				
(³) This	attestation is	only applicable to	Bovidae and Cervidae other than	African buffalo (Syncerus caffer).			
(⁴) This	attestation is	only applicable to	o African buffalo (<i>Syncerus caffer).</i>				
	cination is not d in.	compulsory, but i	the animals have been vaccinated	, information on the vaccine(s) used	and the time of vaccination shall be		
exp	ortation to the	i Únion of the thi	rd country,territory or part thereof		rior to the date of authorisation for or during a period where restrictive part thereof.		

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

		Model S	UI-A			
COL	I.1.		I.2. Certificate reference N		ertificate to El	
t		Address Tel.	I.3. Central competent authority I.4. Local competent authority			
dispatched consignment	1.5.	Consignee Name Address Postal code	1.6.			
ď	1.7.	Tel. Country of origin ISO code I.8. Region of origin Code	I.9. Country of ISO destination	code I.10. Region of destination	Code	
Part I: Details	1.11.	Place of origin	l.12.			
Pai		Name Approval number Address				
		Place of loading Address Approval number	I.14. Date of departure			
	1.15.	Means of transport Aeroplane Ship Railway wagon Road vehicle Other	I.16. Entry BIP in EU			
		Identification Documentary references	1.17.			
	I.18.	Description of commodity	I.19. Commo	odity code (HS code) 01.06.19		
	1.21.			I.20. Quantity	100	
		Seal/Container No		1.24.	103	
		Commodities certified for: Approved body				
	1.26.		I.27. For import or admissio	n into EU		
	1.28.	Identification of the commodities	1			
		Species Identification system (scientific name)	Identification number	Age	Sex	

	COUNT	RY	Model SUI-A							
	II.	Health inf	formation II.a. Certificate reference number II.b.							
	11.1.	Animal h	ealth attestation							
		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.								
ficatio			(a) where the diseases referred to in this certificate are notifiable,							
Certi			(b) which at the date of issuing this certificate has been free for the past 12 months from rinderpest.							
Part II: Certification		II.1.2.	They come from the body, institute or centre/holding (1) described in Box I.11.							
۵.			(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 1.7. 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 <i>Brucella</i> antigen test for porcine brucellosis taken in the 30 days prior to dispatch to the Union.]							

COUNTRY Model SUI-A II.a. Certificate reference number Health information 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.] $(^1)$ or [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.] II.1.8. Classical swine fever (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 classical swine fever.] [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.] (¹) or II.1.9. African swine feve (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 African swine fever.] (¹) or [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, and 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: (¹) [anthrax on the (dd/mm/yyyy) with the following vaccine(s) (name of vaccine (s) used)], (¹) [rabies on the (dd/mm/yyyy) with the following vaccine(s) (name of vaccine (s) used)]. II.1.12. Parasite treatment

	Y				Model SUI-
II. I	Health in	formation		II.a. Certificate reference number	II.b.
I	II.1.13.	described in Box I.	ded for dispatch to the Unior 15. that were cleaned and	n on	ally authorised disinfectant and so
Notes					
				Box I. 28. coming from an approved body, or centre located within a Member State	
Part I:					
— Box r	eference			ntainer and lorries), flight number (aircraft) gnor shall inform the BIP of entry into the	
— Box r	eference	I.28.: Identification s the ISO code	vstem: Specify the identification of the exporting country and	on system (tag, tattoos, brand, chip, trans permit tracing of their premises of origin.	sponder). The identifier shall include
		Age: months.			
		Sex (M = male	e, F = female, C = castrated).	
		Species Select	t the species amongst those	listed below:	
Order		Family	Genera/species		
Artiodact	yla	Suidae	Babyrousa ssp., Hylochoe	erus ssp., Phacochoerus ssp., Potamocho	<i>erus</i> ssp., <i>Sus</i> ssp.
		Tayassuidae	Catagonus ssp., Pecari-Ta	ayassu ssp.	
		Hippopotamidae	Hexaprotodon-Choeropsis	, Hippopotamus ssp.	
Part II:					
(¹) Keep	as appr	opriate.			
(²) Vacci filled		not compulsory, but if	the animals have been vacci	nated, information on the vaccine(s) used	and the time of vaccination must be
	carried 06/2010.		th the protocols that, for the	disease concerned, are described in Pa	art 6 of Annex I to Regulation (EU)
expor	rtation to	the Union of the cour	try, territory or part thereof o	when the animals were loaded either p decribed in Boxes I.7. and I.8., or during a als from that country,territory or part there	a period where restrictive measures
Official v	eterinaria	an			
Name	e (in cap	ital letters):		Qualifica	ation and title:
Date:				Signatur	re:

		Model T	RE-A				
col	JNTR 1.1.	Consignor Name		e reference No	Veterinary c	ertificate to EL	
nent		Address Tel.	1.4. Local competent authority				
dispatched consignment	1.5.	Consignee Name Address	1.6.				
spatched		Postal code Tel.					
đ	1.7.	Country of origin ISO code I.8. Region of origin Code	I.9. Country destinatio		I.10. Region of destination	Code	
Partl : Details	1.11.	Place of origin Name Approval number Address	1.12.				
	I.13.	Place of loading Address Approval number	I.14. Date of departure				
	1.15.	Means of transport Aeroplane Ship Railway wagon Road vehicle Other Locumentary references	I.16. Entry BIP in EU I.17.				
	I.18.	Description of commodity		I.19. Commodity co 0	de (HS code) 1.06.19		
				1.2	0. Quantity		
	1.21.			1.2	2. Number of packa	ges	
		Seal/Container No		1.2	4.		
	1.25.	Commodities certified for: Approved body					
	1.26.		I.27. For impo	ort or admission into	EU 🗌		
	1.28.	Identification of the commodities					
		Species Identification system (scientific name)	Identification	number	Age	Sex	

	COUNTI	RY	Model TRE-4				
	II.	Health inf	ormation II.a. Certificate reference number II.b.				
_	II.1.	Animal health attestation					
		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:					
Part II: Certification		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., 					
Ра			 (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, rables, (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/hold- ing (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.				
		Other vaccinations					
			(a) They have not been vaccinated against rinderpest,				

П.	Health info	ormation		II.a. Certificate reference number	Model TRE		
II.				II.a. Certificate reference number	П.В.		
	(4)	(b) They have beer	n vaccinated against:				
		(¹) [anthrax on the used)],	(dd/mm/yyyy)(d	ate(s)) with the following vaccine(s)	(name of vaccine(s		
 (¹) [rables on the							
						They have been loaded for dispatch to the Union on	
Notes							
				28. coming from an approved body, i r centre located within a Member Sta			
Part I:							
— Box	k reference			er and lorries), flight number (aircraft) r shall inform the BIP of entry into the			
— Box reference I.28.: Identification system: Specify the identification system (tag, tattoos, brand, chip, transponder). The identification system (tag, tattoos, brand, chip, transponder). The identification system is a constrained of the exporting country and permit tracing of their premises of origin.							
		Age: months.					
	Sex (M = male, F = female, C = castrated).						
		Species: Sele	ted below:				
Order		Family	Genera/species				
Perisso	odactyla	Tapiridae	Tapirus ssp.				
		Rhinocerotidae	Ceratotherium ssp., Dicerorhir	nus ssp., Diceros ssp., Rhinoceros ss	p		
Probos	cidea	Elephantidae	Elephas ssp., Loxodonta ssp.				
Part II	:						
(¹) Kee	ep as appro	priate.					
(²) Thi	s attestation	is only applicable to	Rhinocerotidae.				
(³) Thi	s attestation	is only applicable to	<i>Elephas.</i> ssp.				
	ccination is r ed in.	not compulsory, but if	the animals have been vaccinate	d, information on the vaccine(s) used	and the time of vaccination must b		
exp	portation to	the Union of the thi		en the animals were loaded either p f described in Boxes I.7. and I.8., c	or during a period where restrictiv		

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 back-up 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;
 - (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.
- 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.

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.
- 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;
 - (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.'