ANIMAL HEALTH CERTIFICATE FOR THE MOVEMENT BETWEEN MEMBER STATES OF BOVINE ANIMALS NOT INTENDED FOR SLAUGHTER (MODEL 'BOV-INTRA-X')

IRO	PEAN UNI	ION				INTRA
	I.1	Consignor		I.2	IMSOC reference	
		Name		I.2a	Local reference	
		Address		I.3	Central Competent Authority	QR CODE
		Country	ISO country code	I.4	Local Competent Authority	
	1.5	Consignee		I.6	Operator conducting assembly o	operations independently of an
0		Name			establishment Name	Registration No
		Address			Address	
		Country	ISO country code		Country	ISO country code
	I.7	Country of origin	ISO country code	I.9	Country of destination	ISO country code
	I.8	Region of origin	Code	I.10	Region of destination	Code
	I.11	Place of dispatch		I.12	Place of destination	
		Name	Registration/Approval No		Name	Registration/Approval No
		Address			Address	
		Country	ISO country code		Country	ISO country code
	I.13	Place of loading		I.14	Date and time of departure	
	I.15	Means of transport		I.16	Transporter	
		□ Vessel	□ Aircraft		Name	Registration/Authorisation No
					Address	
		Railway	□ Road vehicle		Country	ISO country code
				I.17	Accompanying documents	
		Identification	□ Other		Туре	Code
	Document				Country Commercial document reference	ISO country code
	I.18	Transport conditio	ns 🗆 Ambient		Chilled	Frozen
	I.19	Container number/	Seal number			
		Container No	5	Seal No		

I.20	Certified	as or for									
🗆 Furt	her keeping		□ Slaughter			□ Conf	ined est	ablishment	Germinal	Germinal products	
🗆 Reg	istered equine ani	mal	Travelling cire	cus/anima	l act	Exhibition			□ Event or	□ Event or activity near borders	
🗆 Rele	ease into the wild		Dispatch cents	re		🗆 Relay	ing are	a/purification	Ornamen	tal aquac	ulture
						centre			establishme	ent	
🗆 Furt	□ Further processing □ Organic fertilizers and soil					🗆 Tech	nical us	e	Quarantin	ne or simi	lar
			improvers						establishme	ent	
	lucts for human		D Pollination				•	animals for	□ Other		
consu	mption					human	consum	ption			
I.21	□ For tr	ansit throug	h a third country	У							
	Third co	untry				IS	O count	try code			
	Exit poir	nt				BO	CP code	;			
	Entry po					BC	CP code	•			
I.22	🗆 For transit	through Me	mber State(s)			I.23	⊐ For e	xport			
	Member State		ISO	country co	de	Third country ISO country code			code		
	Member State		ISO	country co	de	Exit point BCP code					
	Member State		ISO	country co	de						
I.24	Estimated journ	ney time				I.25	Jour	ney log	□ yes		🗆 no
I.26	Total number o	f packages				I.27	Total	l quantity			
I.28	Total net weigh	0 0	(D)			I.29	Total	space foreseen	for the cons	ignment	
I.30	Description of c	onsignment									
CN co	ode Spec	eies Subsp	ecies/Category	Sex	Iden syste	tification		Identification r	number	Age	Quantity
					5950						Туре
Region origin		Cold	store		Iden	tification 1	nark	Type of packag	ging		Net weight
Slaug	hterhouse	Treatr	nent type			ure of modity		Number of pac	kages		Batch No
		Date of collect	of tion/production		Man plan	ufacturing t	5	Approval or re number of plant/establish	-	Test	

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II. Heal	th information	1		II.a	Certificate reference	II.b	IMSOC reference		
I, the undersigned official veterinarian, hereby certify									
II.1.	The bovin	e animals ⁽¹⁾	of the consignment desc	cribed in	Part I meet the follow	ing requ	irements:		
	II.1.1.	They are i 2019/2035	dentified as provided f	or in Art	icle 38 of Commissio	n Deleg	ated Regulation (EU)		
	II.1.2.		at least the 30 day period ounger than 30 days of		the departure of the	consign	ment, or since birth, it		
		II.1.2.1.	have been continuous	sly reside	nt in the establishmen	t of orig	in;		
		II.1.2.2.			h kept bovine animal ns for animal health re		ower health status or		
		II.1.2.3.		country	rect contact with kept or territory during the				
II.1.3. They have not shown clin the clinical examination w the consignment, on				as carried	out, within the 24 ho				
II.2.	According to official information, the animals described in Part I meet the following health requirement								
	II.2.1.	They do not come from establishments subject to movement restrictions affecting the specie or situated in a restricted zone established for reasons of diseases listed for bovine animals.							
	II.2.2.	They come from establishments free from infection with <i>Brucella abortus</i> , <i>B. melitensis</i> and <i>B. suis</i> without vaccination regarding bovine animals, and							
	⁽²⁾ either	[the establishments of origin are situated in a Member State or zone thereof with the stat free from infection with <i>Brucella abortus</i> , <i>B. melitensis</i> and <i>B. suis</i> regarding the bovi population;]							
	⁽²⁾ and/or	suis with Delegated during the	be been subjected to a te one of the diagnostic of Regulation (EU) 2020 30 day period prior to sys after parturition;]	methods)/688, car	provided for in Part	l of An ve resul	nex I to Commission ts, on a sample taken		
	⁽²⁾ and/or	[they are 1	ess than 12 months old;]					
	⁽²⁾ and/or	[they are c							
	II.2.3.	They come from establishments free from infection with <i>Mycobacterium tubercula</i> complex (<i>M. bovis, M. caprae</i> and <i>M. tuberculosis</i>), and							
	⁽²⁾ either		ishments of origin are infection with <i>Mycoba</i> <i>is</i>);]						
	⁽²⁾ and/or	(<i>M. bovis,</i> Part 2 of A	been subjected to a to <i>M. caprae</i> and <i>M. tube</i> Annex I to Delegated R 30 day period prior to	erculosis) Regulation	with one of the diagn (EU) 2020/688, carr	ostic m	ethods provided for in		
	⁽²⁾ and/or	[they are]	ess than 6 weeks old.]						

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II.2.4.	They come from establishments in which infection with rabies virus in kept terrestrial animal has not been reported during the 30 day period prior to departure.	ls
П.2.5.	They come from establishments situated in an area of at least 150 km radius around thos establishments in which infection with epizootic haemorrhagic disease virus has not bee reported in kept animals of listed species for that disease during the last 2 years prior t departure.	en
II.2.6.	They come from establishments in which anthrax in ungulates has not been reported durin the 15 days period prior to departure.	g
П.2.7.	They come from establishments in which surra (<i>Trypanosoma evansi</i>) has not been reported during the 30 days period prior to departure, and	d
⁽²⁾ either	[surra has not been reported in the establishments during the last 2 years prior to the departure.]	ir
⁽²⁾ 0r	[surra has been reported during the last 2 years prior to departure, following the last outbreat the affected establishments have remained under movement restrictions until:	k
	- the infected animals have been removed from the establishments, and	
	 the remaining animals on the establishments have been subjected to a test for surr (<i>Trypanosoma evansi</i>) with one of the diagnostic methods provided for in part 3 c Annex I to Delegated Regulation (EU) 2020/688, carried out, with negative results, o samples taken at least 6 months after the infected animals have been removed from the establishments.] 	of m
⁽²⁾ either[II.2.8.	They originate from a Member State or a zone free from infection with bluetongue viru (serotypes 1-24), where no case of infection with bluetongue virus (serotypes 1-24) has bee confirmed during the last 24 months in the targeted animal population and have not bee vaccinated with a live vaccine against infection with bluetongue virus (serotypes 1-24) in th 60 day period before the date of movement and the requirements laid down in Articl $32(1)(a)$, (b) or (c) or Article $32(2)$ of Delegated Regulation (EU) 2020/688 are fulfilled.]	en en ne
⁽²⁾ and/or[II.2.8.	They originate from a Member State or a zone covered by the eradication programme for infection with bluetongue virus (serotypes 1-24) and the requirements laid down in Articl $32(1)(a)$, (b) or (c) or Article $32(2)$ of Delegated Regulation (EU) 2020/688 are fulfilled, an they	le
⁽²⁾ either	[II.2.8.1. have been kept in a Member State or zone seasonally free from infection with bluetongue virus (serotypes 1-24) in accordance with Article 40(3) of Commission Delegated Regulation (EU) 2020/689	
	⁽²⁾ <i>either</i> [II.2.8.1.1. for at least 60 days prior to the date of movement]]	
	(2) and/or [II.2.8.1.2. for at least 28 days prior to the date of movement and have bee subjected to a serological test, with negative results, carried out o samples collected at least 28 days following the entry date of th animal into the Member State or zone seasonally free from infectio with bluetongue virus (serotypes 1-24)]]	n ne
	(2) and/or [II.2.8.1.3. for at least 14 days prior to the date of movement and have bee subjected to a PCR test, with negative results, carried out on sample collected at least 14 days following the entry date of the animal int the Member State or zone seasonally free from infection wit bluetongue virus (serotypes 1-24);]]]	es to

Produced during contingency

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⁽²⁾ and/or	[II.2.8.2.	of destinati	protected against attacks by vectors during transportation to the place ion and have been kept protected against attacks by vectors in a vector stablishment
	⁽²⁾ either	[II.2.8.2.1.	for at least 60 days prior to the date of movement]]
	⁽²⁾ and/or		for at least 28 days prior to the date of movement and have been subjected to a serological test, with negative results, carried out on samples collected at least 28 days following the date of the commencement of the period of protection against attacks by vectors]]
	⁽²⁾ and/or	[11.2.8.2.3.	for at least 14 days prior to the date of movement and have been subjected to a PCR test, with negative results, carried out on samples collected at least 14 days following the date of the commencement of the period of protection against attacks by vectors;]]]
⁽²⁾ and/or	[II.2.8.3.	bluetongue	vaccinated against those serotypes from 1 to 24 of infection with virus which were reported during the past 2 years in that Member ne and are within the immunity period guaranteed in the specifications ine and
	⁽²⁾ either	[11.2.8.3.1.	have been vaccinated more than 60 days before the date of movement]]
	⁽²⁾ and/or	[11.2.8.3.2.	have been vaccinated with an inactivated vaccine and subjected to a PCR test, with negative results on samples collected at least 14 days after the onset of the immunity set in the specifications of the vaccine;]]]
⁽²⁾ and/or	[II.2.8.4.	specific an	subjected with positive results to a serological test able to detect tibodies against all serotypes 1-24 of infection with bluetongue virus uring the past 2 years in that Member State or zone and
	⁽²⁾ either	[II.2.8.4.1.	the serological test has been carried out on samples collected at least 60 days before the date of movement.]]]
	⁽²⁾ and/or	[11.2.8.4.2.	the serological test has been carried out on samples collected at least 30 days before the date of the movement and the animal has been subjected to a PCR test, with negative results, carried out on samples collected not earlier than 14 days before the date of movement.]]]
⁽²⁾ and/or[II.2.8.	virus (ser bluetongue	otypes 1-24) virus (serot	Member State or a zone neither free from infection with bluetongue nor covered by the eradication programme for infection with ypes 1-24) and the requirements laid down in Article 32(1)(a), (b) or belegated Regulation (EU) 2020/688 are fulfilled, and they
⁽²⁾ either	[II.2.8.1.	of destinati	protected against attacks by vectors during transportation to the place on and have been kept protected against attacks by vectors in a vector stablishment
	⁽²⁾ either	[II.2.8.1.1.	for at least 60 days prior to the date of movement]]
	⁽²⁾ and/or		for at least 28 days prior to the date of movement and have been subjected to a serological test, with negative results, carried out on samples collected at least 28 days following the date of the commencement of the period of protection against attacks by vectors]]
	⁽²⁾ and/or	[II.2.8.1.3.	for at least 14 days prior to the date of movement and have been subjected to a PCR test, with negative results, carried out on samples collected at least 14 days following the date of the commencement of the period of protection against attacks by vectors;]]]

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⁽²⁾ and/or [II.2.	situated in establishm Sections	h kept for the 60 day period prior to departure in an establishment a Member State or in an area of at least 150 km radius centred on the nent, where surveillance in compliance with the requirements set out in and 2 of Chapter 1 of Part II of Annex V to Delegated Regulation 0/689 has been carried out during that period, and
(²⁾ eith	er [II.2.8.2.1.	the animals have been vaccinated against those serotypes from 1 to 24 of infection with bluetongue virus which were reported during the past 2 years in an area of at least 150 km radius centred on the place where the animals were kept and are within the immunity period guaranteed in the specifications of the vaccine and
	⁽²⁾ either	[II.2.8.2.1.1. have been vaccinated more than 60 days before the date of movement]]]
	⁽²⁾ and/or	[II.2.8.2.1.2. have been vaccinated with an inactivated vaccine and subjected to a PCR test, with negative results on samples collected at least 14 days after the onset of the immunity set in the specifications of the vaccine;]]]]
⁽²⁾ ana	/or [II.2.8.2.2.	the animals have been immunised against those serotypes from 1 to 24 of infection with bluetongue virus which were reported during the past 2 years in an area of at least 150 km radius centred on the place where the animals were kept, and
	⁽²⁾ either	[II.2.8.2.2.1. the animals have been subjected with positive results to a serological test carried out on samples collected at least 60 days before the date of movement]]]
	⁽²⁾ and/or	[II.2.8.2.2.2. the animals have been subjected with positive results to a serological test carried out on samples collected at least 30 days before the date of the movement and to a PCR test, with negative results, carried out on samples collected not earlier than 14 days before the date of movement.]]]]
Part I the M	I of Annex V to	requirements laid down in points 1 to 3 of Section 1 of Chapter 2 of Delegated Regulation (EU) 2020/689 and the competent authority of rigin authorised movement of those animals to another Member State
⁽²⁾ either [II.2.	Member Member	tatus free from infection with bluetongue virus (serotypes 1-24) and the State of destination has informed the Commission and the other States that such movement is authorised subject to the conditions o in Article $43(2)(a)$, (b) and (c) of Delegated Regulation (EU) and
⁽²⁾ eith	er [II.2.8.1.1.	point 5 of Section 1 of Chapter 2 of Part II of Annex V to that Delegated Regulation, and
⁽²⁾ ana	or [II.2.8.1.2.	point 6 of Section 1 of Chapter 2 of Part II of Annex V to that Delegated Regulation, and

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(-	²⁾ and/or		point 7 of Section 1 of Chapter 2 of Part II of Annex V to that Delegated Regulation, and
6	²⁾ and/or		point 8 of Section 1 of Chapter 2 of Part II of Annex V to that Delegated Regulation, and
			the requirements laid down in Article 32(1)(a), (b) or (c) or Article 32(2) of Delegated Regulation (EU) 2020/688 and the requirements laid down in Article 33 of that Delegated Regulation are fulfilled.]]]
⁽²⁾ and/or [[11.2.8.2.	(serotypes Commission subject to th	proved eradication program for infection with bluetongue virus $1-24$) and the Member State of destination has informed the n and the other Member States that such movement is authorised ne conditions referred to in Article $43(2)(a)$, (b) and (c) of Delegated (EU) 2020/689 and
6	²⁾ either		point 5 of Section 1 of Chapter 2 of Part II of Annex V to that Delegated Regulation, and
(²⁾ and/or		point 6 of Section 1 of Chapter 2 of Part II of Annex V to that Delegated Regulation, and
6	²⁾ and/or	-	point 7 of Section 1 of Chapter 2 of Part II of Annex V to that Delegated Regulation, and
(-	²⁾ and/or	[11.2.8.2.4.	point 8 of Section 1 of Chapter 2 of Part II of Annex V to that Delegated Regulation, and
			the requirements laid down in Article 32(1)(a), (b) or (c) or Article 32(2) of Delegated Regulation (EU) 2020/688 and the requirements laid down in Article 33 of that Delegated Regulation are fulfilled.]]]
⁽²⁾ and/or [[11.2.8.3.	the eradicat and the Mer	from infection with bluetongue virus (serotypes 1-24) nor covered by ion programme for infection with bluetongue virus (serotypes 1-24) mber State of destination has informed the Commission and the other ates that such movement is authorised
(-	²⁾ either	[11.2.8.3.1.	without any conditions, and
6	²⁾ and/or	-	subject to the conditions referred to in point 5 of Section 1 of Chapter 2 of Part II of Annex V to Delegated Regulation (EU) 2020/689, and
(.	²⁾ and/or	-	subject to the conditions referred to in point 6 of Section 1 of Chapter 2 of Part II of Annex V to Delegated Regulation (EU) $2020/689$, and
6	²⁾ and/or		subject to the conditions referred to in point 7 of Section 1 of Chapter 2 of Part II of Annex V to Delegated Regulation (EU) 2020/689, and
6	²⁾ and/or	-	subject to the conditions referred to in point 8 of Section 1 of Chapter 2 of Part II of Annex V to Delegated Regulation (EU) 2020/689, and
			the requirements laid down in Article 32(1)(a), (b) or (c) or Article 32(2) of Delegated Regulation (EU) 2020/688 and the requirements laid down in Article 33 of that Delegated Regulation are fulfilled.]]]

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(2)[(2)either[II	.2.9. They	are moved t	to a Member State or zone thereof with the status free from enzootic
	bovine leul	kosis, and	
⁽²⁾ either	[II.2.9.1.	they come	from establishments free from enzootic bovine leukosis.]]
⁽²⁾ or	[II.2.9.1.	enzootic bo	from establishments not free from enzootic bovine leukosis, and ovine leukosis has not been reported in those establishments during the period prior to departure, and
	⁽²⁾ either	[II.2.9.1.1.	they are over 24 months of age and they have been subjected to a serological test for enzootic bovine leukosis with one of the diagnostic methods provided for in Part 4 of Annex I to Delegated Regulation (EU) 2020/688, carried out with negative results
		⁽²⁾ either	[II.2.9.1.1.1. on samples taken on two occasions at an interval of at least four months while kept in isolation from the other bovine animals of the establishment]]]
	⁽²⁾ and/or	⁽²⁾ and/or	[II.2.9.1.1.2. on a sample taken during the 30 day period prior to the departure of the consignment, and all bovine animals over 24 months of age kept in the establishment have been subjected to a serological test for enzootic bovine leukosis with one of the diagnostic methods provided for in Part 4 of Annex I to Delegated Regulation (EU) 2020/688, carried out, with negative results, on samples taken on two occasions at an interval of not less than four months during the 12 month period prior to the departure of the consignment;]]]] they are less than 24 months of age and they were horn to dam.
	⁽²⁾ and/or	[11.2.9.1.2.	they are less than 24 months of age and they were born to dam subjected to a serological test for enzootic bovine leukosis with one of the diagnostic methods provided for in Part 4 of Annex I to Delegated Regulation (EU) 2020/688, carried out, with negative results, on samples taken on two occasions at an interval of not less than four months during the 12 month period prior to the departure of the consignment.]]]
⁽²⁾ or[II.2.9.	They are n for enzooti	noved to a M	ember State or zone thereof with an approved eradication programme cosis, and
⁽²⁾ either	[II.2.9.1.		from establishments free from enzootic bovine leukosis.]]
⁽²⁾ or	[II.2.9.1.	they come enzootic bo	from establishments not free from enzootic bovine leukosis, and ovine leukosis has not been reported in those establishments during the period prior to departure, and
	⁽²⁾ either	[II.2.9.1.1.	they are over 24 months of age and they have been subjected to a serological test for enzootic bovine leukosis with one of the diagnostic methods provided for in Part 4 of Annex I to Delegated Regulation (EU) 2020/688, carried out with negative results

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		⁽²⁾ either	[II.2.9.1.1.1.	on samples taken on two occasions at an interval of at least four months while kept in isolation from the other bovine animals of the establishment]]]
		⁽²⁾ and/or	[Ш.2.9.1.1.2.	on a sample taken during the 30 day period prior to the departure of the consignment, and all bovine animals over 24 months of age kept in the establishment have been subjected to a serological test for enzootic bovine leukosis with one of the diagnostic methods provided for in Part 4 of Annex I to Delegated Regulation (EU) 2020/688, carried out, with negative results, on samples taken on two occasions at an interval of not less than four months during the 12 month period prior to the departure of the consignment;]]]]
	⁽²⁾ and/or	[II.2.9.1.2.	which has be leukosis with Annex I to I negative resu not less than	than 24 months of age and they were born to dam, een subjected to a serological test for enzootic bovine one of the diagnostic methods provided for in Part 4 of Delegated Regulation (EU) 2020/688, carried out, with lts, on samples taken on two occasions at an interval of four months during the 12 month period prior to the he consignment.]]]]
⁽²⁾ [⁽²⁾ either[infectious l	bovine rhino	tracheitis/infe	Member State or zone thereof with the status free from ctious pustular vulvovaginitis and they have not been ninotracheitis/infectious pustular vulvovaginitis, and
⁽²⁾ either	[II.2.10.1.			establishments free from infectious bovine pustular vulvovaginitis, and
	⁽²⁾ either	[II.2.10.1.1	thereof wit	nents of origin are situated in a Member State or zone the status free from infectious bovine s/infectious pustular vulvovaginitis]]
	⁽²⁾ and/or	[II.2.10.1.2	prior to depa the detection (BoHV-1) wi of Annex I to result, carried	have been subjected to quarantine for at least 30 days rture and have been subjected to a serological test for a of antibodies against whole bovine herpes virus-1 th one of the diagnostic methods provided for in Part 5 to Delegated Regulation (EU) 2020/688, with a negative out on a sample taken during the 15 day period prior to of the consignment.]]]
(2) ₀ r	[II.2.10.1.	rhinotrache approved o have been whole BoH Annex I to	itis/infectious quarantine esta subjected to a IV-1, with on Delegated Re sample taken	tablishments not free from infectious bovine pustular vulvovaginitis and they have been kept in an ablishment for at least 30 days prior to departure and serological test for the detection of antibodies against e of the diagnostic methods provided for in Part 5 of gulation (EU) 2020/688, with a negative result, carried not less than 21 days after commencement of the

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	hey are moved to a Member State or zone thereof with an approved eradication nme for infectious bovine rhinotracheitis/infectious pustular vulvovaginitis, and
⁽²⁾ either [II.2.10	
⁽²⁾ eithei	 [II.2.10.1.1. the establishments of origin are situated in a Member State or zone thereof with the status free from infectious bovine rhinotracheitis/infectious pustular vulvovaginitis]]
⁽²⁾ and/o	<i>The stablishments of origin are situated in a Member State or zone thereof with an approved eradication programme for infectious bovine rhinotracheitis/infectious pustular vulvovaginitis]</i>
⁽²⁾ and/c	or [II.2.10.1.3. the animals have been subjected to quarantine for at least 30 days prior to departure and have been subjected to a serological test for the detection of antibodies against whole bovine herpes virus-1 (BoHV-1) with one of the diagnostic methods provided for in Part 5 of Annex I to Delegated Regulation (EU) 2020/688 with a negative result, carried out on a sample taken during the 15 day period prior to the departure of the consignment]]
⁽²⁾ and/c	or [II.2.10.1.4. the animals are destined for an establishment which keeps bovine animals for meat production without contact to bovine animals of other establishments, and from which they are directly moved to the slaughterhouse.]]]
⁽²⁾ or [II.2.10	0.1. they come from establishments not free from infectious bovine rhinotracheitis/infectious pustular vulvovaginitis, and
	 they have been kept in an approved quarantine establishment for at least 30 days prior to departure, and
	 they have been subjected to a serological test for the detection of antibodies against whole BoHV-1, with one of the diagnostic methods provided for in Part 5 of Annex I to Delegated Regulation (EU) 2020/688, with a negative result, carried out on a sample taken not less than 21 days after commencement of the quarantine.]]]
⁽²⁾ [⁽²⁾ <i>either</i> [II.2.11.	They are moved to a Member State or zone thereof with the status free from viral diarrhoea and they have not been vaccinated against bovine viral diarrhoea, and
⁽²⁾ either [II.2.1] ⁽²⁾ either	.1. they come from establishments free from bovine viral diarrhoea, and
⁽²⁾ and/c	as referred in point 1(c)(ii) or (iii) of Section 2 of Chapter 1 of Part VI of Annex IV to Delegated Regulation (EU) 2020/689, carried out, with negative results, within the four months period prior to the departure of the consignment]]
⁽²⁾ and/c	<i>or</i> [II.2.11.1.3. the animals have been tested individually to exclude the presence of bovine viral diarrhoea virus prior to the departure of the consignment.]]]

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(2) ₀ r	[II.2.11.1.	they come from establishments not free from bovine viral diarrhoea and they have been subjected to a test for bovine viral diarrhoea virus antigen or genome with one of the diagnostic methods provided for in Part 6 of Annex I to Delegated Regulation (EU) 2020/688, carried out with negative results, and
	⁽²⁾ either	[II.2.11.1.1. they have been kept in an approved quarantine establishment for a period of at least 21 days prior to the departure of the consignment
		⁽²⁾ [and in case of pregnant dams, they have been subjected to a serological test for the detection of antibodies against bovine viral diarrhoea virus with one of the diagnostic methods provided for in Part 6 of Annex I to Delegated Regulation (EU) 2020/688, carried out, with negative results, on samples taken not less than 21 days after commencement of the quarantine]]]
	⁽²⁾ and/or	[II.2.11.1.2. they have been subjected to a serological test for the detection of antibodies against bovine viral diarrhoea virus with one of the diagnostic methods provided for in Part 6 of Annex I to Delegated Regulation (EU) 2020/688, with positive results,
		⁽²⁾ either [II.2.11.1.2.1. in case of non-pregnant animals, carried out on samples taken prior to departure of the consignment]]]
		⁽²⁾ and/or [II.2.11.1.2.1. in case of pregnant dams, carried out on samples taken before insemination preceding the current gestation.]]]]
⁽²⁾ or[II.2.11		loved to a Member State or zone thereof with an approved eradication programme viral diarrhoea, and
⁽²⁾ either	[II.2.11.1.	
	⁽²⁾ either	[II.2.11.1.1. the establishments of origin are situated in a Member State or zone thereof with the status free from bovine viral diarrhoea]]
	⁽²⁾ and/or	[II.2.11.1.2. the establishments of origin are situated in a Member State or zone thereof with an approved eradication programme for bovine viral diarrhoea]]
	⁽²⁾ and/or	[II.2.11.1.3. the establishments of origin have been subjected to a testing regime as referred in point 1(c)(ii) or (iii) of Section 2 of Chapter 1 of Part VI of Annex IV to Delegated Regulation (EU) 2020/689, carried out, with negative results, within the last 4 months prior to the departure of the consignment]]
	⁽²⁾ and/or	[II.2.11.1.4. the animals have been tested individually to exclude the presence of bovine viral diarrhoea virus prior to the departure of the consignment]]
	⁽²⁾ and/or	[II.2.11.1.5. the animals are destined for an establishment which keeps bovine animals for meat production separate from bovine animals of other establishments, and from which they are directly moved to the slaughterhouse]]

EUROPEAN UNION		Certificate model BOV-INTRA-X	
⁽²⁾ and/or	-	they come from establishments not free from bovine viral diarrhoea and they have been subjected to a test for bovine viral diarrhoea virus antigen or genome with one of the diagnostic methods provided for in Part 6 of Annex I to Delegated Regulation (EU) 2020/688, carried out with negative results, and	
	⁽²⁾ either	[II.2.11.2.1. they have been kept in an approved quarantine establishment for a period of at least 21 days prior to the departure of the consignment ⁽²⁾ [and in case of pregnant dams, they have been subjected to a serological test for the detection of antibodies against bovine viral diarrhoea virus with one of the diagnostic methods provided for in Part 6 of Annex I to Delegated Regulation (EU) 2020/688, carried out, with negative results, on samples taken not less than 21 days	
		after commencement of the quarantine]]]	
	⁽²⁾ and/or	[II.2.11.2.2. they have been subjected to a serological test for the detection of antibodies against bovine viral diarrhoea virus with one of the diagnostic methods provided for in Part 6 of Annex I to Delegated Regulation (EU) 2020/688, with positive results,	
		⁽²⁾ <i>either</i> [II.2.11.2.2.1. in case of non-pregnant animals, carried out on samples taken prior to departure of the consignment]]]	
		(2) and/or [II.2.11.2.2.1. in case of pregnant dams, carried out on samples taken before insemination preceding the current gestation.]]]]]	
		owledge and as declared by the operator, the animals come from establishments onormal mortalities with an undetermined cause.	
²⁾ [II.4. According to official information and as declared by the operator, they are semen			
II.4.1.		e from a semen collection centre and will be transported directly to another semen a centre in accordance with Article 19 of Commission Delegated Regulation (EU) ; and	
⁽²⁾ either [II.4.2.			
⁽²⁾ or [II.4.2. they were a Chapter I of admission t quarantine a II.4.3. the prior co been obtain		subjected, with negative results, to all tests referred to in point 1(b) and (c) of of Part 1 of Annex II to Delegated Regulation (EU) 2020/686, required before to a semen collection centre carried out during the period immediately preceding and during the quarantine period; and] onsent of the centre veterinarian of the semen collection centre of destination has need by the operator; and	
		II.4.4.	the means

PEAN UNION	Certificate model BOV-INTRA-
an	is certificate is valid for 10 days from the date of issuing. In the case of transport by waterway/sea of imals, the period of validity of the certificate may be extended by the duration of the journey betterway/sea.
as	nce leaving their establishments of origin and before arriving to this establishment approved for sembly operations, none of the animals of the consignment has undergone more than two assemble erations, and
⁽²⁾ eit	<i>her</i> [they come from their establishments of origin.]]
⁽²⁾ or	[at least one of the animals of the consignment has undergone one assembly operation on a approved establishment.]]
⁽²⁾ or	[at least one of the animals of the consignment has undergone two assembly operations o approved establishments.]]
Animal we	Ifare attestation
with the	of inspection, the animals covered by this health certificate were fit to be transported in accordance provisions of Council Regulation (EC) No $1/2005$ on the intended journey due to stat (insert date) ⁽⁴⁾⁽⁵⁾ .
Notes:	
from the E Protocol on	the with the Agreement on the withdrawal of the United Kingdom of Great Britain and Northern Irelan uropean Union and the European Atomic Energy Community, and in particular Article 5(4) of the Ireland / Northern Ireland in conjunction with Annex 2 to that Protocol, references to European Unio ficate include the United Kingdom in respect of Northern Ireland.
	l health certificate shall be completed according to the notes for the completion of certificates provide

Part I:

Box reference I.11:	<i>"Place of dispatch":</i> Indicate an establishment of the origin of the animals in the consignment or an establishment approved for assembly operations in accordance with Articles 97 and 99 of Regulation (EU) 2016/429 of the European Parliament and of the Council.
Box reference I.12:	<i>"Place of destination":</i> Indicate an establishment of the final destination of the consignment or an establishment approved for assembly operations in accordance with Articles 97 and 99 of Regulation (EU) 2016/429.
Box reference I.17:	"Accompanying documents": In case the animals are dispatched from an establishment approved for assembly operations in the Member State of origin, the reference number(s) of the official document(s), based on which the animal health certificate for this consignment is issued in this establishment approved for assembly operations, may be indicated.
	In case the animals are dispatched from an establishment approved for assembly operations in the Member State of passage, the reference number(s) of the certificate(s), based on which the animal health certificate for this consignment is issued in this establishment approved for assembly operations, must be indicated.
Box reference I.30:	<i>"Identification number"</i> : Indicate identification codes of the animals in the consignment identified in accordance with Article 38 of Delegated Regulation (EU) 2019/2035.

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Part II:						
(1)	⁽¹⁾ There can be one or more animals in the consignment.					
(2)	Delete if not applicable.					
(3)	Applicable in case the consignment is dispatched fi operations.	rom the establishment approved for assembly				
(4)	In the case where a consignment is grouped in an establishment approved for assembly operations and comprises animals that were loaded on different dates, the date which the journey commenced for the whole consignment is considered to be the earliest date when any part of the consignment left the establishment of origin.					
(5)	This statement does not exempt transporters from their obligation in accordance with Union rules in force in particular regarding the fitness to be transported.					
Official	veterinarian					
Name (in	capital letters)	Qualification and title				
Local Co	ntrol Unit name	Local Control Unit code				
Date						
Stamp		Signature				