I.1. Consignor					I.2. IMSOC reference	I.2.a. Local refere	ence		
	Name					I.3. Central Comp	etent Authority		
	Address Country		ISO Code		I.4. Local Competent Autho				
t									
en	I.5. Consignee				I.6. Operator conducting assemb establishment	bly operations independ	dently of an		
H	Name Address				Name				
igi	Country		ISO Code		Address				
ns	country		100 0040		Approval Number				
CO					Country	ISO Code			
of									
ion	I.7. Country of ori	igin		ISO Code	I.9. Country of destination		ISO Code		
Part I: Description of consignment	I.8. Region of orig	in		Code	I.10. Region of destination		Code		
esc	I.11. Place of disp	atch			I.12. Place of destination				
Õ	Name	aten			Name				
t I:	Address				Address				
ar	Approval				Approval Number				
Ч	Nûmber		100 0-1-			ICO Cada			
	Country		ISO Code		Country	ISO Code			
	I.13. Place of load	ing			I.14. Date and time of departure				
	Name								
	Address								
	Approval Number								
	Country		ISO Code						
	I.15. Means of Tra	ansport			I.16. Transporter				
	Mode	International	Identification		Name				
		transport document			Address				
					Approval Number				
					Country	ISO Code			
					<ul> <li>I.17. Accompanying documents</li> <li>Document Type</li> <li>Accompanying document reference</li> <li>Date of Issue</li> </ul>				
					Country				
	140 Turner out of				Place of issue				
	I.18. Transport co Ambient 🗖	nations	Chille		Frozen 🗆				
			Chine	:u 🗀	F102en				
		umber / Seal Numb	ber						
	I.20. Certified as			-	<b>N</b> 1 1 1 1 1				
	Other 🗆		Further keeping		Exhibition	Release into the v Quarantine or sir			
	Confined establis	hment 🗀	Travelling circus/a	animal act 🗀	Breeding 🗆	establishment			
	Event or activity	near borders 🛛							
		rough a third cou	ntry						
	Third country				ISO Code				
	Exit point Entry point				BCP code BCP code				
		nrough Member St	ate(s)		I.23. For export				
		a subrimerimer st							
	Member State		ISO Code		Third country Exit point	ISO Code BCP code			
	I.24. Estimated jo	urney time			I.25. Journey Log	Der coue			
	I.27. Total quantit				I.28. Total gross weight				
	I.30. Description of				_				
	1. 01 LIVE ANIMA	-							
		-							

# en

### INTRA

0102 Live bovine animals								
E	Bison spp							
#1.	Commodity	Subcategory	Sex	Identification system				
Species		Identification Number	Age	Quantity				

	II. Health info	rmation								
	I, the under	rsigned offic	cial veterina	arian, hereby certify tha	L t:					
	II.1.									
	II.1.1.	They are identified as provided for in Article 38 of Commission Delegated Regulation (EU) 2019/2035.								
on	II.1.2.	-	t least 30 da an 30 days		eparture of the consignment	or since birth, if they are				
cati	II.1.2.1.	have been continuously resident in the establishment of origin;								
Part II: Certification	II.1.2.2.	have not been in contact with kept bovine animals of a lower health status or subject to movement restrictions for animal health reasons;								
t II: C	II.1.2.3.	have not been in direct or indirect contact with kept animals that have entered the Union from a third country or territory during the last 30 days prior to the date of departure of the animals.								
Par	II.1.3.	They have not shown clinical signs or symptoms of diseases listed for bovine animals during the clinical examination which was carried out, within the last 24 hours prior to the time of departure of the consignment, on (insert date dd/mm/yyyy).								
	II.2.	According requireme		formation, the animals	described in Part I meet the	following health				
	II.2.1.	(2)	○ either	affecting bovine anima species or diseases sub	-	ns of diseases listed for those				
	<ul> <li>(2) or [They come from establishments or zones subject to movement restrictions affecting bov animals and established for (3), but derogations from movement restriction have been granted, and:</li> </ul>									
		□ (2)	[they comp	bly with the requirement	ts set out in (4)	;]]				
		□ (2)	[and in par	rticular, they are	(5).]]					
	II.2.2.	-		lishments free from infe egarding bovine animals	ection with Brucella abortus, , and:	B. melitensis and B. suis				
	(2) 🗆 either			•	Member State or zone thereo and B. suis regarding the bov					
	(2) 🗆 and/or	of the diag 2020/688, c	nostic meth arried out,	ods provided for in Part with negative results, on	n with Brucella abortus, B. m 1 of Annex I to Commission a a sample taken during the l t females taken at least 30 da	ast 30 days prior to the date				
	(2) 🗆 and/or	[they are le	ess than 12 i	months old;]						
	(2) 🗆 and/or	[they are ca	astrated.]							
	II.2.3.	-		lishments free from infe I. tuberculosis), and:	ection with Mycobacterium to	ıberculosis complex (M.				
	(2) □ either			-	Member State or zone therec mplex (M. bovis, M. caprae a					
	<ul> <li>(2) □</li> <li>(2) □</li> <li>(2) □</li> <li>(3) □</li> <li>(4) □</li> <li>(5) □</li> <li>(6) □</li> <li>(7) □</li> <li>(7) □</li> <li>(8) □</li> <li>(9) □</li> <li>(10) □</li> <li>(10</li></ul>									
	(2) 🗆 and/or	[they are le	ess than 6 w	eeks old.]						
	II.2.4.	-			-	ot terrestrial animals has not consignment.				
	II.2.5.	<ul> <li>been reported during the last 30 days prior to the date of departure of the consignment.</li> <li>2.5. They come from establishments situated in an area of at least 150 km radius around those establishments in which infection with epizootic haemorrhagic disease virus:</li> </ul>								

	II. Health infor	rmation							
	(2) □ either	[has not been reported in kept animals of listed species for that disease during the last 2 years prior to the date of departure of the consignment;]							
	(2) 🗆 and/or	[has been reported in kept animals of listed species for that disease during the last 2 years prior to date of departure of the consignment and the animals have been kept in a zone seasonally free fir epizootic haemorrhagic disease in accordance with Parts 1 and 2 of Annex IX to Delegated Regula (EU) 2020/688:							
		(2)							
		(2) 🗆 and/or	for at least 28 days prior to the dat subjected to a serological test, with 28 days following the date of entry	negative results, carried out	on samples collected at least				
Par		(2) 🗆 and/or	for at least 14 days prior to the dat subjected to a PCR test, with negati days following the date of entry of	ve results, carried out on sam	ples collected at least 14				
	(2) 🗆 and/or	[has been reported in kept animals of listed species for that disease during the last 2 years prior to the date of departure of the consignment and the animals have been protected against attacks by vectors during transportation to the place of destination and they have been kept protected against attacks by vectors in a vector protected establishment fulfilling the requirements provided for in Part 3 of Anno IX to Delegated Regulation (EU) 2020/688:							
		(2) 🗆 either	for at least 60 days prior to the dat	e of departure of the consign	nent;]				
		(2) 🗆 and/or	for at least 28 days prior to the dat subjected to a serological test, with 28 days following the date of the co by vectors;]	negative results, carried out	on samples collected at least				
		(2) 🗆 and/or	for at least 14 days prior to the dat subjected to a PCR test, with negati days following the date of commer vectors;]	ve results, carried out on sam	ples collected at least 14				
	(2) □ and/or	date of dep	reported in kept animals of listed sp parture of the consignment and the n and the other Member States that	has informed the					
	II.2.6.	-	from establishments in which anth to the date of departure of the const		reported during the last 15				
	II.2.7.		from establishments in which surr s prior to the date of departure of th		not been reported during the				
	(2) 0 either	[surra has departure.]	not been reported in the establishm ]	ents during the last 2 years p	rior to the date of their				
	(2) or	[surra has been reported during the last 2 years prior to the date of departure of the consignment, and following the date of the last outbreak, the affected establishments have remained under movement restrictions until the date on which the infected animals have been removed from the establishments, and the remaining animals in the establishments have been subjected to a test for surra with one of the diagnostic methods provided for in Part 3 of Annex I to Delegated Regulation (EU) 2020/688, carried out, with negative results, on samples taken at least 6 months following the date on which the infected animals have been removed from the establishments.]							
	(2) □ either	[II.2.8.	They originate from a Member Sta virus (serotypes 1-24), where no ca been confirmed in the targeted and date of departure of the consignment against infection with bluetongue of departure of the consignment and (b) or (c), or Article 32(2) of Delega	se of infection with bluetong mal population during the las ent, and have not been vaccin virus (serotypes 1-24) in the la nd the requirements laid dow	ue virus (serotypes 1-24) has st 24 months prior to the ated with a live vaccine ast 60 days prior to the date n in Article 32(1), points (a),				
	(2) 🗆 and/or	[II.2.8.	They originate from a Member Sta programme for infection with blue						

-	IUN				2024/1044 (2021	(403) Model BOV-INTRA-2			
	II. Health info	rmation							
				rticle 32(1), points (a), (b re fulfilled, and they:	) or (c), or Article 32(2) of De	legated Regulation (EU)			
		(2) 🗆 either	[II.2.8.1.	with bluetongue virus	ember State or zone thereof s (serotypes 1-24) in accordan Regulation (EU) 2020/689:	seasonally free from infectior ce with Article 40(3) of			
Caro	(2) 🗆 either	[II.2.8.1.1.	for at leas	t 60 days prior to the date	e of depature of the consign	ment;]]			
	(2) 🗆 and/or	[II.2.8.1.2.	subjected 28 days fo	for at least 28 days prior to the date of departure of the consignment and have been subjected to a serological test, with negative results, carried out on samples collected at leas 28 days following the date of entry of the animals into the Member State or zone thereof seasonally free from infection with bluetongue virus (serotypes 1-24);]]					
TMT	(2) 🗆 and/or	[II.2.8.1.3.	subjected days follow	for at least 14 days prior to the date of departure of the consignment and have been subjected to a PCR test, with negative results, carried out on samples collected at least 14 days following the date of entry of the animals into the Member State or zone thereof seasonally free from infection with bluetongue virus (serotypes 1-24);]]]					
		(2) 🗆 and/or	[II.2.8.2.			ing transportation to the against attacks by vectors in a			
	(2) 🗆 either	[II.2.8.2.1.	for at leas	t 60 days prior to the dat	e of departure of the consigr	nment;]]			
	(2) 🗆 and/or	[II.2.8.2.2.	subjected	to a serological test, with llowing the date of the co		nment and have been t on samples collected at leas of protection against attacks			
	(2) 🗆 [II.2.8.2.3. and/or		subjected	to a PCR test, with negati wing the date of the com	e of departure of the consigr ve results, carried out on sam mencement of the period of				
		(2) □ and/or	[II.2.8.3.	bluetongue virus whicl during the last 2 years					
	(2) 🗆 either	[II.2.8.3.1.	have been	vaccinated more than 6	0 days prior to the date of de	eparture of the consignment;]			
	(2) □ and/or	[11.2.8.3.2.	results, ca		lected at least 14 days after t	ed to a PCR test, with negative he date of the onset of the			
		(2) □ and/or	[II.2.8.4.	specific antibodies again reported in that Memb	ith positive results to a serol inst all serotypes 1-24 of infe er State or zone thereof duri f the consignment, and:	ection with bluetongue virus			
	(2) 🗆 either	[II.2.8.4.1.		gical test has been carrie parture of the consignme	d out on samples collected a ent.]]]	t least 60 days prior to the			
	(2) □ [II.2.8.4.2. the serological test has been carr and/or date of departure of the consign with negative results, carried ou date of departure of the consign			parture of the consignme tive results, carried out o	ent and the animals have bee on samples collected not earl	en subjected to a PCR test,			
	(2) 🗆 and/or [II.2.8.	(serotypes (serotypes	1-24) nor co 1-24) and th	overed by the eradication	n programme for infection w vn in Article 32(1), points (a)	fection with bluetongue virus vith bluetongue virus ), (b) or (c), or Article 32(2) of			
		(2) 🗆 either	[II.2.8.1.		ainst attacks by vectors dur d have been kept protected a				

	NION 2024/1044 (2021/403) WOULD BOV-INTRA-						
	II. Health infor	rmation					
				vector protected establ	ishment:		
	(2) 🗆 either	[II.2.8.1.1.	for at least	-	e of departure of the consigni	ment;]]	
	(2) 🗆 and/or	[II.2.8.1.2.	subjected t	o a serological test, with lowing the date of the co	e of departure of the consign negative results, carried out ommencement of the period o	on samples collected at least	
<u>Part II: Ceruncauon</u>	(2) 🗆 and/or	[II.2.8.1.3.	subjected t	o a PCR test, with negati	e of departure of the consigni ve results, carried out on sam mencement of the period of p	ples collected at least 14	
Par		(2) 🗆 and/or	[II.2.8.2.	consignment in an esta of at least 150 km radiu compliance with the re	last 60 days prior to the date blishment situated in a Mem is centred on the establishme quirements set out in Part II, ted Regulation (EU) 2020/689	ber State or within an area nt, where surveillance in Chapter 1, Sections 1 and	
	(2) 🗆 either	[II.2.8.2.1.	bluetongue of the cons	e virus which were repo ignment within an area ere kept and are within	against all serotypes from 1 to rted during the last 2 years pr of at least 150 km radius cen the immunity period guarant	rior to the date of departure tred on the place where the	
		(2) 🗆 either	[II.2.8.2.1. 1.	have been vaccinated r consignment;]]]	nore than 60 days prior to the	e date of departure of the	
		(2) □ and/or	[II.2.8.2.1. 2.	with negative results of	vith an inactivated vaccine an n samples collected at least 14 the specifications of the vacci	4 days after the date of onset	
	(2) 🗆 and/or	[II.2.8.2.2.	the animals have been immunised against all serotypes from 1 to 24 of infection with bluetongue virus which were reported during the last 2 years prior to the date of departure of the consignment 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.		subjected with positive resulted at least 60 days prior to the	ts to a serological test carried date of departure of the	
		(2) □ and/or	[II.2.8.2.2. 2.	out on samples collecte consignment and to a F	subjected with positive result d at least 30 days prior to the PCR test, with negative results an 14 days prior to the date of	, carried out on samples	
	<ul> <li>(2) □ and/ They do not fulfil the requirements laid down or[II.2.8. Delegated Regulation (EU) 2020/689 and the construction authorised movement of those animals to ano</li> </ul>			EU) 2020/689 and the co	mpetent authority of the Mer	nber State of origin	
		(2) 🗆 either	[II.2.8.1.	the Member State of de Member States that suc	m infection with bluetongue estination has informed the Co ch movement is authorised su B(2), points (a), (b) and (c), of l	ommission and the other bject to the conditions	
	(2) □ either	[II.2.8.1.1.	Part II, Cha	pter 2, Section 1, point 5	5, of Annex V to that Delegate	d Regulation, and	
	(2) 🗆 and/or	[II.2.8.1.2.	Part II, Cha	pter 2, Section 1, point 6	6, of Annex V to that Delegate	d Regulation, and	
and/or (2) [II.2.8.1.3. Part II, Chapter 2, Section 1, point 7, of Ar and/or			pter 2, Section 1, point 7	, of Annex V to that Delegate	d Regulation, and		

	II. Health info	rmation					
	(2) 🗆 and/or	[II.2.8.1.4.	Part II, Cha	pter 2, Section 1, point 8	8, of Annex V to that Delegated	d Regulation, and	
					or (c), or Article 32(2) of Dele that Delegated Regulation are		
Part II: Certification		(2) 🗆 and/or	[II.2.8.2.	(serotypes 1-24) and th Commission and the ot	lication programme for infect e Member State of destination ther Member States that such ns referred to in Article 43(2), EU) 2020/689, and:	n has informed the movement is authorised	
	(2) □ either	[II.2.8.2.1.	Part II, Cha	pter 2, Section 1, point s	5, of Annex V to that Delegated	d Regulation, and	
Par	(2) □ and/or	[II.2.8.2.2.	Part II, Cha	pter 2, Section 1, point (	6, of Annex V to that Delegated	d Regulation, and	
	(2) 🗆 and/or	[II.2.8.2.3.	Part II, Cha	pter 2, Section 1, point '	7, of Annex V to that Delegated	d Regulation, and	
	(2) 🗆 and/or	[II.2.8.2.4.	Part II, Cha	pter 2, Section 1, point 8	8, of Annex V to that Delegated	d Regulation, and	
					or (c), or Article 32(2) of Deleg that Delegated Regulation are		
		(2) 🗆 and/or	[II.2.8.3.	by the eradication prog 24) and the Member St	ction with bluetongue virus (se gramme for infection with blu ate of destination has informe hat such movement is authori	letongue virus (serotypes 1- ed the Commission and the	
	(2) □ either	[II.2.8.3.1.	without an	y conditions, and:			
	(2) 🗆 and/or	[II.2.8.3.2.	subject to the conditions referred to in Part II, Chapter 2, Section 1, point 5, of Annex V to Delegated Regulation (EU) 2020/689, and				
	(2) 🗆 and/or	[II.2.8.3.3.	subject to the conditions referred to in Part II, Chapter 2, Section 1, point 6, of Annex V to Delegated Regulation (EU) 2020/689, and				
	(2) 🗆 and/or	[II.2.8.3.4.		he conditions referred t Regulation (EU) 2020/68	to in Part II, Chapter 2, Sectior 9, and	n 1, point 7, of Annex V to	
	(2) 🗆 and/or	[II.2.8.3.5.		he conditions referred t Regulation (EU) 2020/68	to in Part II, Chapter 2, Sectior 9, and	n 1, point 8, of Annex V to	
	the requirements laid down in Article 32(1), points (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) [(2)   0 either	[II.2.9.	They are m leukosis, ai		e or zone thereof with the stat	us free from enzootic bovine	
		(2) o either	[II.2.9.1.	they come from establi	ishments free from enzootic b	ovine leukosis.]]]	
		(2) or	[II.2.9.1.	enzootic bovine leukos	ishments not free from enzoo sis has not been reported in th or to the date of departure of t	ose establishments during	
	(2) □ either	[II.2.9.1.1.	enzootic bo	ovine leukosis with one	d they have been subjected to of the diagnostic methods pro D/688, carried out with negativ	wided for in Part 4 of Annex	
		(2) 🗆 either	[II.2.9.1.1. 1.	-	vo occasions at an interval of ther bovine animals of the est	-	
		(2) 🗆 and/or	[II.2.9.1.1. 2.	consignment, and all b establishment have be leukosis with one of th	ing the last 30 days prior to th ovine animals over 24 month en subjected to a serological t e diagnostic methods provide EU) 2020/688, carried out, wit	s of age kept in the est for enzootic bovine d for in Part 4 of Annex I to	

	1011				(		
	II. Health info	rmation					
					occasions at an interval of not r to the date of departure of t		
	(2) 🗆 and/or	[II.2.9.1.2.	test for enz Annex I to samples ta	zootic bovine leukosis w Delegated Regulation (E ken on two occasions at	e and they were born to dams ith one of the diagnostic meth (U) 2020/688, carried out, with an interval of not less than 4 ure of the consignment.]]]	ods provided for in Part 4 of negative results, on	
ertuno	(2) or	[II.2.9.	They are moved to a Member State or zone thereof with an approved eradication programme for enzootic bovine leukosis, and:				
		(2) ○ either	[II.2.9.1.	they come from establi	shments free from enzootic b	ovine leukosis.]]	
Part		(2) or	[II.2.9.1.	enzootic bovine leukos	shments not free from enzoo is has not been reported in th r to the date of departure the	ose establishments during	
	(2) □ either	[II.2.9.1.1.	enzootic bo	ovine leukosis with one	d they have been subjected to of the diagnostic methods pro D/688, carried out with negativ	wided for in Part 4 of Annex	
		(2) □ either	[II.2.9.1.1. 1.		vo occasions at an interval of her bovine animals of the est		
		(2) 🗆 and/or	[II.2.9.1.1. 2.	consignment, and all b establishment have bee leukosis with one of th Delegated Regulation ( samples taken on two o	ing the last 30 days prior to th ovine animals over 24 months en subjected to a serological to e diagnostic methods provide EU) 2020/688, carried out, wit occasions at an interval of not or to the date of departure of t	s of age kept in the est for enzootic bovine d for in Part 4 of Annex I to h negative results, on less than 4 months during	
	(2) 🗆 and/or	[II.2.9.1.2.	they are less than 24 months of age and they were born to dams, 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 4 months during the last 12 months prior to the date of departure of the consignment.]]]]				
	(2) [(2) 0 either	[II.2.10.	bovine rhi	notracheitis/infectious p	e or zone thereof with the stat oustular vulvovaginitis and th cheitis/infectious pustular vul	ey have not been vaccinated	
		(2) 0 either	[II.2.10.1.		shments free from infectious ous pustular vulvovaginitis, ar		
	(2) □ either	[II.2.10.1.1			uated in a Member State or zo racheitis/infectious pustular v		
	(2) 🗆 and/or	[II.2.10.1.2	departure of antibodi methods pi negative re	of the consignment and ies against whole bovine rovided for in Part 5 of A	o quarantine for at least 30 da have been subjected to a sero e herpes virus-1 (BoHV-1) with Annex I to Delegated Regulation mple taken during the last 15	logical test for the detection one of the diagnostic on (EU) 2020/688, with a	
		(2) or	[II.2.10.1.	rhinotracheitis/infection approved quarantine endeparture of the consignation of antiboon the detection of antiboon methods provided for in with a negative result,	shments not free from infections pustular vulvovaginitis an establishment for at least 30 da gnment and have been subjection dies against whole BoHV-1, with an Part 5 of Annex I to Delegat carried out on a sample taker nent of the quarantine.]]]	d they have been kept in an ays prior to the date of ted to a serological test for ith one of the diagnostic ed Regulation (EU) 2020/688,	
	(2) or	[II.2.10.			e or zone thereof with an appr hinotracheitis/infectious pust		

	II. Health info	rmation						
		(2) o either	[II.2.10.1. they come from establishments free from infectious bovine rhinotracheitis/infectious pustular vulvovaginitis, and:					
	(2) 🗆 either	[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]]]]					
Certification	(2) □ and/or	[II.2.10.1.2	the establishments of origin are situated in a Member State or zone thereof with an approved eradication programme for infectious bovine rhinotracheitis/infectious pustular vulvovaginitis]]]]					
Part II: Certifi	(2) 🗆 and/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 last 15 days prior to the date of departure of the consignment]]]]					
д	(2) 🗆 and/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.2.10.1. they come from establishments not free from infectious bovine rhinotracheitis/infectious pustular vulvovaginitis, and:					
	_	-	been kept in an approved quarantine establishment for at least 30 days prior to the date of of the consignment, and					
	_	one of the o with a nega	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 the date of commencement of the quarantine.]]]					
	(2) [(2) 0 either	[II.2.11.	They are moved to a Member State or zone thereof with the status free from bovine viral diarrhoea and they have not been vaccinated against bovine viral diarrhoea, and:					
		(2) ○ either	[II.2.11.1. they come from establishments free from bovine viral diarrhoea, and:					
	(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 have been subjected to a testing regime as referred in Part VI, Chapter 1, Section 2, point 1(c)(ii) or (iii), of Annex IV to Delegated Regulation (EU) 2020/689, carried out, with negative results, within the 4 months period prior to the date of departure of the consignment,]]]]					
	(2) 🗆 and/or	[II.2.11.1.3	the animals have been tested individually to exclude the presence of bovine viral diarrhoea virus prior to the date of departure of the consignment.]]]]					
		(2) or	[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 date of departure of the consignment, [and in the 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 the date of commencement of the quarantine;] (2) ]]					
	(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 in the case of non-pregnant dams, carried out on samples taken prior to the date.1.of departure of the consignment]]]]					
		(2) 🗆 and/or	[II.2.11.1.2 in the case of pregnant dams, carried out on samples taken before the date of.1.insemination preceding the current gestation.]]]]					
	(2) or	[II.2.11.	They are moved to a Member State or zone thereof with an approved eradication					

	II. Health info	rmation							
			programm	e for bovine viral diarrh	loea, and:				
		(2) 🗆 either			shments free from bovi	ne vir	al diarrhoea, and:		
uo	(2) 🗆 either	[II.2.11.1.1		shments of origin are sit povine viral diarrhoea;]]		e or zo	one thereof with the status		
ificati	(2) 🗆 and/or	•	approved e	eradication programme	uated in a Member Stat for bovine viral diarrho	oea;]]]			
irt II: Certi	(2) □ and/or (2) □ and/or (2) □ and/or	[II.2.11.1.3	Chapter 1, s	the establishments of origin have been subjected to a testing regime as referred in Part VI, Chapter 1, Section 2, point 1(c)(ii) or (iii), of Annex IV to Delegated Regulation (EU) 2020/689, carried out, with negative results, within the last 4 months prior to the date of departure of the consignment;]]]					
Pa	(2) 🗆 and/or	[II.2.11.1.4		s have been tested indiv to the date of departure		resen	ce of bovine viral diarrhoea		
	(2) 🗆 and/or	[II.2.11.1.5	production	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;]]]					
		(2) □ and/or	[II.2.11.2.	have been subjected to with one of the diagnos		liarrh or in F			
	(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 date of departure of the consignment, [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 the date of commencement of the quarantine;] (2)]						
	(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) □ either	[II.2.11.2.2 .1.	in the case of non-preg of departure of the con		on sar	nples taken prior to the date		
		(2) 🗆 and/or	[II.2.11.2.2 .1.		dams, carried out on sa g the current gestation.]	-	s taken before the date of		
	II.3.		-	•	by the operator, the anir th an undetermined cau		ome from establishments		
	II.4.		ents are mac (EU) 2020/6		gnment in accordance v	with A	rticle 4 of Delegated		
	II.5.	This animal health certificate is valid for 10 days from the date of issuing. In the case of transport by waterway/sea of animals, the period of validity of the certificate may be extended by the duration of the journey by waterway/sea.							
	(2)(6) 🗆 [II.6.	establishm	ent approve		ons, none of the animals		he date of arrival to this e consignment has		
	(2) ○ either	[they come	from their	establishments of origir	ı.]]				
	(2) or	[at least on establishm		mals of the consignmen	t has undergone one ass	embly	operation in an approved		
	(2) or	[at least on establishm		mals of the consignmen	t has undergone two ass	sembly	y operations in the approved		
	Animal we	lfare attesta	tion						
		-		-	nal health certificate we No 1/2005 on the intend		-		

	II. Health info	rmation							
		(insert date) (7) (8).							
	Notes:	(insert date) (7) (6).							
	In accorda from the Eu Protocol or	In accordance with the Agreement on the withdrawal of the United Kingdom of Great Britain and Northern Ireland from the European Union and the European Atomic Energy Community, and in particular Article 5(4) of the Protocol on Ireland/Northern Ireland in conjunction with Annex 2 to that Protocol, references to the Union in this animal health certificate include the United Kingdom in respect of Northern Ireland.							
rtifica	This animal health certificate shall be completed in accordance with the notes for the completion of certificates provided for in Chapter 2 of Annex I to Commission Implementing Regulation (EU) 2020/2235.								
Ce	Part I:								
art	Box reference I.11:	"Place of dispatch": Indicate an establishment establishment approved for assembly operation (EU) 2016/429 of the European Parliament and	ons in accordance with Article						
	Box reference I.12:	"Place of destination": Indicate an establishme establishment approved for assembly operation (EU) 2016/429.							
	Box reference I.17:	"Accompanying documents": In case the animals are dispatched from an establishment approved for							
		In case the animals are dispatched in the Member State of passage, th which the animal health certificate approved for assembly operations	e reference number(s) of the e for this consignment is issue	certificate(s), based on					
	Box reference I.30:	"Identification number": Indicate identification codes of the animals in the consignment identified in							
	Part II:								
	(1)	There may be one or more animals in the cons	signment.						
	(2)	Delete if not applicable.							
	(3)	Insert the name of the disease(s).							
	(4)	Insert the specific reference to the article(s), ti Commission providing for those requirements		nt legal act(s) adopted by the					
	(5)	Insert the specific attestation(s) provided for in Commission, as referred to in Article 126(1), pe							
	(6)	Applicable in case the consignment is dispatch operations.	ed from the establishment ap	proved for assembly					
	(7)	In the case where a consignment is grouped in comprises animals that were loaded on differed whole consignment is considered to be the ear establishment of origin.	ent dates, the date which the j	ourney commenced for the					
	(8)	This statement does not exempt transporters f force in particular regarding the fitness to be t		dance with Union rules in					
	Certifying Offi Name (in capi	icer/Official veterinarian tal letters)	Qualification and title						
	Date		Signature						
	Stamp								