EUROPEAN UNION INTRA

	I.1. Consignor					I.2. IMSOC ref	erence	I.2.a. Lo	cal reference	
	Name								tral Competen	t Authority
	Address								al Competent	
	Country ISO Code								•	,
	,									
Ħ	I.5. Consignee					I.6. Operator of establishment	conducting assembly of	perations	independent	ly of an
of consignment	Name									
뒫	Address					Address	Name			
٠	Country			ISO Code		Approval Nu	mhar			
ű							Country ISO Code			
ၓ						Country				
ð	I.7. Country of ori	gin			ISO Code	I.9. Country of	f destination			ISO Code
on										
ΡĖ	I.8. Region of original	in			Code	I.10. Region of	f destination			Code
Part I: Description	I.11. Place of dispa					I.12. Place of o				
esc	Name					Name				
À	Address					Address				
ţ	Approval Numbe	r				Approval Nu	mber			
ar	Country			ISO Code		Country			ISO Code	
д	-									
	I.13. Place of loadi	ing				I.14. Date and	time of departure			
	Name									
	Address									
	Approval Numbe	r								
	Country			ISO Code						
	I.15. Means of Tra	nsport				I.16. Transpor	ter			
	Mode	Internation	nal	Identification		Name				
	Mode	transport	iiui	lucitimeuton		Address				
		document				Activity ID				
						Country		ISO	Code	
						_	nying documents			
						Commercial document Date of issue				
						reference Country Place of issue				
	I.18. Transport co	nditions						133410		
	I.19. Container No									
	1,10, 001,011,01	,, 0001110								
	I.20. Certified as									
	Event or activity r	near borders	\Box	Slaughter \square		Quarantine or similar Other Cestablishment Exhibition Exhibition				
	Confined establish	amont \square		Travelling circus/an	imal act \square					
	Release into the w			Travelling circus/an	Illiai act 🗀					
	Release Into the W	пи Ш								
	I.21. For transit th	rough a thir	d coun	try						
	Third country					ISO Code				
	Exit point					BCP code	BCP code			
	Entry point					BCP code				
	I.22. For transit th	rough Mem	ber Sta			I.23. For export				
	Member State			ISO Code		Third country ISO Code				
						Exit point			BCP code	
						I.25. Journey Log I.28. Total gross weight				
	I.27. Total quantit	У				1.28. Total gro	ss weight			
	I.30. Description o	of consignme	ent							
	Commodity	- 3	Specie	98	Sex		Identification system	1	Identification	Number
	Sommoury		Specie		0011		- activities of the state of th	,		
	Ouantitu					Age	<u>'</u>			
	Quantity					Age				

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animals have been removed from the establishments.]

	KOPLAN (
	II. Health info	ormation							
tion	(2)	either □ [II.2.8.	virus (sero been confi the date of against inf the date of	types 1-24), rmed in the departure of ection with departure of	where no ca targeted ani of the consign bluetongue v of the consign	te or a zone thereof free from se of infection with bluetongs mal population during the penment and have not been vacyirus (serotypes 1-24) during soment and the requirements of Delegated Regulation (EU)	ne virus (serotypes 1-24) has briod of 24 months prior to cinated with a live vaccine the period of 60 days prior to laid down in Article 32(1),		
Part II: Certification	(2)	and/or □ [II.2.8.	programm down in A	e for infecti	on with blue points (a), (b)	te or a zone thereof covered k tongue virus (serotypes 1-24) or (c), or Article 32(2) of Delo	and the requirements laid		
Part	(2)		either □ [II.2.8.1.	with bluet	kept in a Member State or zone thereof seasonally free from infection ongue virus (serotypes 1-24) in accordance with Article 40(3) of on Delegated Regulation (EU) 2020/689:				
	(2)			either □ [II.2.8.1.1.	_	od of at least 60 days prior to nt;]]	the date of departure of the		
	(2)			and/or □ [II.2.8.1.2.	consignment negative re- of at least 2 Member Sta	od of at least 28 days prior to nt and have been subjected to sults, carried out on samples 8 days following the date of e ate or zone thereof seasonally virus (serotypes 1-24)]]	a serological test, with collected during the period ntry of the animal into the		
	(2)			and/or □ [II.2.8.1.3.	consignment results, carr 14 days foll State or zor	od of at least 14 days prior to nt and have been subjected to ried out on samples collected owing the date of entry of the ne thereof seasonally free from types 1-24);]]]	a PCR test, with negative during the period of at least animal into the Member		
	(2)		and/or □ [II.2.8.2.	place of de		ainst attacks by vectors durir d have been kept protected ag ishment:			
	(2)			either □ [II.2.8.2.1.	for the peri	od of at least 60 days prior to nt;]]	the date of departure of the		
	(2)			and/or □ [II.2.8.2.2.	consignment negative re- of at least 2	od of at least 28 days prior to nt and have been subjected to sults, carried out on samples 8 days following the date of the cotection against attacks by ve	a serological test, with collected during the period he commencement of the		
	(2)			and/or □ [II.2.8.2.3.	consignment results, care 14 days foll	od of at least 14 days prior to nt and have been subjected to ried out on samples collected owing the date of the comme against attacks by vectors;]]]	a PCR test, with negative during the period of at least		
	(2)		and/or □ [II.2.8.3.	bluetongue Member S	e virus which tate or zone t thin the peri	gainst those serotypes from 1 n were reported during the pe hereof prior to the date of de od of immunity guaranteed i	eriod of 2 years in that parture of the consignment		
	(2)			either □ [II.2.8.3.1.		vaccinated during the period of departure of the consignm			
	(2)			and/or □ [II.2.8.3.2.	have been we PCR test, with during the J	vaccinated with an inactivate th negative results, carried or period of at least 14 days afte et in the specifications of the	d vaccine and subjected to a ut on samples collected r the date of the onset of the		

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	II. Health info	rmation					
	(2)		and/or □ [II.2.8.4.	specific an reported in	tibodies aga n that Memb	rith positive results to a serologinst all serotypes 1-24 of infector State or zone thereof during of the consignment, and:	ction with bluetongue virus
ıtion	(2)			either □ [II.2.8.4.1.		cical test has been carried out of at least 60 days prior to the nt;]]	
Part II: Certification	(2)			and/or □ [II.2.8.4.2.	the period consignme negative re	gical test has been carried out of at least 30 days prior to the nt and the animal has been so esults, carried out on samples ter than 14 days prior to the d nt;]]]	date of departure of the abjected to a PCR test, with collected during the period
	(2)	and/or □ [II.2.8.	bluetongue with bluet	ree from infection with n programme for infection d down in Article 32(1),) 2020/688 are fulfilled, and			
	(2)		either □ [II.2.8.1.	place of de		gainst attacks by vectors durin Id have been kept protected a lishment:	
	(2)			either □ [II.2.8.1.1.		iod of at least 60 days prior to nt;]]	the date of departure of the
	(2)			and/or □ [II.2.8.1.2.	consignme negative re of at least 2	iod of at least 28 days prior to nt and have been subjected to esults, carried out on samples 28 days following the date of to protection against attacks by v	o a serological test, with collected during the period he commencement of the
	(2)			and/or □ [II.2.8.1.3.	consignme results, car 14 days fol	iod of at least 14 days prior to nt and have been subjected to rried out on samples collected lowing the date of the comme against attacks by vectors;]]]	a PCR test, with negative during the period of at least
	(2)		and/or □ [II.2.8.2.	consignme least 150 k complianc	ent in an esta m radius cen e with the re I to Delegate	period of 60 days prior to the ablishment situated in a Mem ntred on the establishment, w equirements set out in Part II, ed Regulation (EU) 2020/689 ha	ber State or in an area of at here surveillance in Chapter 1, Sections 1 and 2,
	(2)			either □ [II.2.8.2.1.	24 of infect the period consignme place wher	s have been vaccinated again tion with bluetongue virus wh of 2 years prior to the date of nt in an area of at least 150 km re the animals were kept and guaranteed in the specificatio	nich were reported during departure of the m radius centred on the are within the period of
	(2)				either □ [II.2.8.2.1. 1.	have been vaccinated during days prior to the date of dep	g the period of more than 60 arture of the consignment;]]]
	(2)				and/or □ [II.2.8.2.1. 2.	have been vaccinated with a subjected to a PCR test, with on samples collected during after the date of the onset of specifications of the vaccine	negative results, carried out the period of at least 14 days the immunity set in the

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	II. Health info	rmation					
	(2)			and/or □ [II.2.8.2.2.	24 of infect the period consignment	s have been immunised again ion with bluetongue virus wh of the past 2 years prior to the nt in an area of at least 150 kr e the animals were kept, and:	nich were reported during e date of departure of the n radius centred on the
ification	(2) either [II.2.8.2.2. 1.					results to a serological test ca orior to the date of departure	
Part II: Certification	(2)			and/or □ [II.2.8.2.2. 2.	test carried days prior t test, with n the period	s have been subjected with po out on samples collected dur to the date of departure of the egative results, carried out or of not earlier than 14 days pri gnment.]]]]	ring the period of at least 30 e consignment and to a PCR a samples collected during
	(2)	and/or □ [II.2.8.	Annex V to	Delegated :	Regulation (I	s laid down in Part II, Chapter EU) 2020/689 and the compete nt of those animals to anothe	ent authority of the Member
	(2)		either □ [II.2.8.1.	the Member S	er State of de tates that suc o in Article 43	m infection with bluetongue s estination has informed the Co ch movement is authorised su B(2), points (a), (b) and (c), of I	ommission and the other bject to the conditions
	(2)			either □ [II.2.8.1.1.		pter 2, Section 1, point 5, of A]]	nnex V to that Delegated
	(2)			and/or □ [II.2.8.1.2.		pter 2, Section 1, point 6, of A]]	nnex V to that Delegated
	(2)			and/or □ [II.2.8.1.3.		pter 2, Section 1, point 7, of A]]	nnex V to that Delegated
	(2)			and/or □ [II.2.8.1.4.	Regulation, (a), (b) or (cand the recaption)	pter 2, Section 1, point 8, of A and the requirements laid do c), or Article 32(2) of Delegated quirements laid down in Artic (EU) 2020/688 are fulfilled;]]]	own in Article 32(1), points d Regulation (EU) 2020/688
	(2)		and/or □ [II.2.8.2.	(serotypes Commission subject to	1-24) and the on and the ot the condition	ication programme for infect e Member State of destination her Member States that such as referred to in Article 43(2), EU) 2020/689 and in	n has informed the movement is authorised
	(2)			either \square [II.2.8.2.1.		pter 2, Section 1, point 5, of A]]	nnex V to that Delegated
	(2)			and/or \square [II.2.8.2.2.		pter 2, Section 1, point 6, of A]]	nnex V to that Delegated
	(2)			and/or □ [II.2.8.2.3.		pter 2, Section 1, point 7, of A]]	nnex V to that Delegated
	(2)			and/or □ [II.2.8.2.4.	Regulation, (a), (b) or (c and the reg	pter 2, Section 1, point 8, of A and the requirements laid d c), or Article 32(2) of Delegated uirements laid down in Artic 88 are fulfilled;]]]	own in Article 32(1), points d Regulation (EU) 2020/688
	(2)		and/or □ [II.2.8.3.	by the erac 24) and the	dication prog e Member St	tion with bluetongue virus (so gramme for infection with blu ate of destination has informo nat such movement is authori	netongue virus (serotypes 1- ed the Commission and the

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]	EU	ROPEAN UNION			2023/308 (2021/	403) MODEL CER-INTRA-X
		II. Health information				
		(2)	either □ [II.2.8.3.1.	without any	y conditions;]] and	
		(2)	and/or □ [II.2.8.3.2.	•	he conditions referred to in P Annex V to Delegated Regulat	-
	ion	(2)	and/or □ [II.2.8.3.3.	•	he conditions referred to in P Annex V to Delegated Regulat	-
	tificat	(2)	and/or □ [II.2.8.3.4.	,	he conditions referred to in P Annex V to Delegated Regulat	•
	Part II: Certification	(2)	and/or □ [II.2.8.3.5.	point 8, of A requirement Article 32(2 requirement	he conditions referred to in P Annex V to Delegated Regulat nts laid down in Article 32(1), 2) of Delegated Regulation (EU nts laid down in Article 33 of re fulfilled.]]]	ion (EU) 2020/689, and the points (a), (b) or (c), or () 2020/688 and the
		II.2.9. V	Vith regard to chronic	wasting dis	ease (CWD), they:	
			(2) either °	[II.2.9.1.	are moved from a Member S State listed in Chapter A, Sec VIII to Regulation (EU) 999/20 Parliament and of the Counc	tion C, point 1.1, of Annex 001 of the European
			(2) or ○	[II.2.9.2.	are semi-domesticated reind an area in Finland listed in C 1.2(a), of Annex VIII to Regul seasonal grazing in Finland.]	Chapter A, Section C, point ation (EU) 999/2001 for
			(2) or ○	[II.2.9.3.	are semi-domesticated reind an area in Sweden listed in C 1.2(b), of Annex VIII to Regul seasonal grazing in Norway, sporting or cultural events in grazing in Sweden, or for spo Sweden, and the competent a given its prior written conse	Chapter A, Section C, point lation (EU) 999/2001 after or after having taken part in a Norway, or for seasonal orting or cultural events in authority of Sweden has
			(2) or ○	[II.2.9.4.	are semi-domesticated reind Norway in the area located be Finnish border, and the Norway Fence and are returning to F	oetween the Norwegian- wegian-Finnish Reindeer
			(2) or ○	[II.2.9.5.	are moved from an area in M Norway with a transit throug the competent authority of S its prior written consent to s	gh Sweden or Finland, and weden or Finland has given
			(2) or ∘	[II.2.9.6.	are moved from an area in S Section C, point 1.2(b), of And 999/2001 to Norway, and the Norway has given its prior w movement.]	nex VIII to Regulation (EU) competent authority of
			(2) or ∘	[II.2.9.7.	are forest reindeer moved fr in Chapter A, Section C, point Regulation (EU) 999/2001 to l authority of Finland has give to such movement.]	t 1.2(b), of Annex VIII to Finland, and the competent

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	TOI LIN	UNION			2025/506 (2021/405) MODEL CER-INTRA-A	
	II. Health inf	formation				
cation			(2) or ○	[II.2.9.8.	are moved from an area in a Member State listed in Chapter A, Section C, point 1.1, of Annex VIII to Regulation (EU) 999/2001, other than an area listed in Chapter A, Section C, point 1.2, of Annex VIII to that Regulation, to another Member State listed in Chapter A, Section C, point 1.1, of Annex VIII to that Regulation, or to Norway, and the competent authority of destination has given its prior written consent to such movement.]	
Part II: Certification		το.	(2) or o	[II.2.9.9.	are moved from a confined establishment, as defined in Article 4(48) of Regulation (EU) 2016/429, in a Member State listed in Chapter A, Section C, point 1.1, of Annex VIII to Regulation (EU) 999/2001, to a confined establishment, as defined in Article 4(48) of Regulation (EU) 2016/429, in another Member State, and the competent authority of the Member State of destination has given its prior written consent to such movement.]	
		(2) III.2.10.	infectious approved rhinotrac come from rhinotrac	s bovine rhir eradication heitis/infecti n an establis heitis/infecti during the p	Member State or zone thereof with the status free from otracheitis/infectious pustular vulvovaginitis or with an programme for infectious bovine ous pustular vulvovaginitis in bovine animals and they thment in which infectious bovine ous pustular vulvovaginitis in cervid animals has not been eriod of 30 days prior to the date of departure of the	
	II.3.				by the operator, the animals come from establishments ith an undetermined cause.	
	II.4.	Arrangements are ma Regulation (EU) 2020,	-	port the cons	signment in accordance with Article 4 of Delegated	
	II.5.		nals, the per		ays from the date of issuing. In the case of transport by ty of the certificate may be extended by the duration of the	
	(2) (3) \square [II.6.		ed for asser	mbly operati	of origin and prior to the date of arrival to this ons, none of the animals of the consignment has ns, and:	
	(2)	either \circ [they come f	rom their es	tablishments of origin.]]	
	(2)		ast one of throwed estab		f the consignment has undergone one assembly operation	
	(2)			ne animals o olishments.]]	f the consignment has undergone two assembly operations	
	Animal welfare attestation					
	At the time of inspection, the animals covered by this animal health certificate were fit to be transported in accordance with the provisions of Council Regulation (EC) No 1/2005 on the intended journey due to start on (insert date).					

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EUROPEAN UNION 2023/308 (2021/403) MODEL CER-INTRA-X II. Health information Notes: 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 European Union in this animal health certificate include the United Kingdom in respect of Northern Ireland. Certification 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. Part I: ä Box "Place of dispatch": Indicate an establishment of the origin of the animals in the consignment or an refer I.11: reference 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. "Place of destination": Indicate an establishment of the final destination of the consignment or an Box reference establishment approved for assembly operations in accordance with Articles 97 and 99 of Regulation (EU) 2016/429. I.12: Box "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), reference based on which the animal health certificate for this consignment is issued in this establishment I.17: 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. "Identification number": Indicate identification codes of the animals in the consignment identified in Box reference accordance with Article 73 or Article 74 of Delegated Regulation (EU) 2019/2035. 1.30: Part II: (1) There can be one or more animals in the consignment. (2) Delete if not applicable. (3) Applicable in case the consignment is dispatched from the establishment approved for assembly operations. Certifying Officer/Official veterinarian Name (in capital letters) Authority name Date of signature Signature Stamp

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