EUROPEAN UNION INTRA

	I.1. Consignor				I.2. IMSOC reference	I.2.a. Local referen	ce		
	Name Address					I.3. Central Compet	ent Authority		
	Country		ISO Code			I.4. Local Competer	nt Authority		
of consignment	I.5. Consignee Name Address Country		ISO Code		I.6. Operator conducting assembly operations independently of an establishment Name Address Approval Number				
[co					Country	ISO Code			
ion of			ISC	O Code	I.9. Country of destination		ISO Code		
ipti	I.8. Region of origi		Co	de	I.10. Region of destination		Code		
Part I: Description	I.11. Place of dispa Name Address Approval Number Country		ISO Code		I.12. Place of destination Name Address Approval Number Country	ISO Code			
	I.13. Place of loadi	ing			I.14. Date and time of departure				
	Name Address Approval Number Country		ISO Code		,				
	I.15. Means of Tra	nsport			I.16. Transporter				
	Mode	International transport	Identification		Name				
		document			Address Activity ID				
					Country	ISO Code			
					I.17. Accompanying documents				
					Accompanying document reference Date of issue Country Place of issue				
	I.18. Transport con	nditions	_	_	Frozen 🗆				
	Ambient \square		Chilled [
	I.19. Container No	/ Seal No							
	I.20. Certified as Travelling circus/a	animal act \square	Quarantine or similar establishment \square	c	Organic fertilizers and soil improvers □	Confined establish	ment \square		
	Slaughter \square		Dispatch centre \square		Further processing \square	Registered equidae			
	Germinal products		Exhibition \square Further keeping \square		Event or activity near borders Ornamental aquaculture establishment	Release into the wi	ld 🗆		
	□ Relaying □		Live aquatic animals consumption \Box	for human	Technical use □				
	I.21. For transit th	rough a third coun	ntry						
	Third country Exit point				ISO Code BCP code				
	Entry point				BCP code				
		rough Member Sta			I.23. For export	[]			
	Member State		ISO Code		Third country Exit point	ISO Code BCP code			
	I.24. Estimated jou				I.25. Journey Log				
	I.27. Total quantity	-			I.28. Total gross weight				
	I.30. Description o 1. 01 LIVE ANIMA	-							
	T. OI LIVE AMIMA	ш							

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EUROPEAN UNION INTRA

	Weighing more than 185 g 010599 Other 01059910 Ducks									
	#1. Commodity	Subcategory	Identification system	Identification Number						
	Species	Age	Quantity							
Part I: Description of consignment		лде	Quantity							

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]	II. Health info	ormation									
]	I, the unde	ersigned offi	icial veterin	arian, herek	by certify, tha	ıt:					
]	II.1.	.The animals(1) in the consignment described in Part I meet the following requirements:									
		II.1.1.		ïned establi ion (EU) 201		spatch is approved in accord	ance with A	rticles 97 and 99			
CCI (IIICa II OII		II.1.2.	listed in A	nnex of Con on, or wher 48 hour per	nmission Imp e this is not p	or symptoms of diseases, in plementing Regulation (EU) 2 possible, a clinical inspection departure of the consignmen	018/1882, du , which was	ring the clinical			
•	II.2.		to official in uirements:	nformation,	animals in tl	ne consignment described in	Part I meet	the following			
۱		II.2.1.	•		ifined establi to be moved.	shment that is not subject to	movement 1	restrictions			
	(2)(3) 🗆 either	[II.2.2.	(serotypes confirmed vaccinated 60 day per	They originate from a Member State or a zone free from infection with bluetongue virus (serotypes 1-24), where no case of infection with bluetongue virus (serotypes 1-24) has been confirmed during the last 24 months in the targeted animal population and have not been vaccinated with a live vaccine against infection with bluetongue virus (serotypes 1-24) in the 60 day period before the date of movement and the requirements laid down in Article 32(1)(a), (b) or (c) or Article 32(2) of Commission Delegated Regulation (EU) 2020/688 are							
- 1	(2)(3) □ and/or	[II.2.2.	infection v	with blueton	igue virus (se	te or a zone covered by the e erotypes 1-24) and the requir f Delegated Regulation (EU) 2	ements laid	down in Article			
	(2)	□ either	[II.2.2.1.	2.2.1. 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)	\square either	[II.2.2.1.1.	for at least	60 days prior to the date of n	novement]]				
		(2)	□ and/or	[II.2.2.1.2.	subjected to samples col animal into	28 days prior to the date of not a serological test, with negalected at least 28 days follow the Member State or zone songue virus (serotypes 1-24)]	ntive results, ving the entr easonally fre	carried out on y date of the			
		(2)	□ and/or	[II.2.2.1.3.	subjected to collected at the Membe	14 days prior to the date of note a PCR test, with negative releast 14 days following the ear State or zone seasonally frewirus (serotypes 1-24;]]]	sults, carried entry date of	d out on samples the animal into			
((2)	□ and/or	[II.2.2.2.	place of de		ainst attacks by the vectors of the cattacks by the vectors of the cattacks are the cattacks and the cattacks are the cattacks.					
		(2)	\square either	[II.2.2.2.1.	for at least	60 days prior to the date of n	novement]] .				
		(2)	□ and/or	[II.2.2.2.2.	subjected to samples col	28 days prior to the date of notes a serological test, with negalected at least 28 days follow ment of the period of protect	ntive results, ving the date	carried out on of the			
		(2)	□ and/or	[II.2.2.2.3.	subjected to collected at	14 days prior to the date of not a PCR test, with negative re least 14 days following the coff protection against attacks	sults, carried late of the co	d out on samples ommencement of			
((2)	□ and/or	[II.2.2.3.			gainst those serotypes from n were reported during the p					

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	II. Health inf	ormation							
				0	,				
				State or zone and are within the immunity period guaranteed in the specifications of the vaccine and					
		(2)	□ either	[II.2.2.3.1.	have been went]	vaccinated more than 60 da]	ays before the date of		
Part II: Certification		(2)	□ and/or	[II.2.2.3.2.	PCR test, wi		ated vaccine and subjected to a ples collected at least 14 days the specifications of the		
t II: Cert	(2)	□ and/or	[II.2.2.4.	have been subjected with positive results to a serological test able to detect specific antibodies against all serotypes 1-24 of infection with bluetongue virus reported during the past 2 years in that Member State or zone and					
Pal	3	(2)	\square either	[II.2.2.4.1.	_	cal test has been carried or ore the date of movement]	ut on samples collected at least]		
		(2)	□ and/or	[II.2.2.4.2.	[II.2.2.4.2. the serological test has been carried out on samples collected at 30 days before the date of the movement and the animal has been subjected to a PCR test, with negative results, carried out on sam collected not earlier than 14 days before the date of movement;]				
	(2)(3) [II.2.2. They originate from a Member State or a zone neither free from infection with blu virus (serotypes 1-24) nor covered by the eradication programme for infection with bluetongue virus (serotypes 1-24) and the requirements laid down in Article 32(1)(c) (c) or Article 32(2) of Delegated Regulation (EU) 2020/688 are fulfilled, and they								
	(2)	□ either	[II.2.2.1.	have been protected against attacks by the vectors during transportation to the place of destination and have been kept protected against attacks by vectors in a vector protected establishment and					
		(2)	\square either	[II.2.2.1.1.	for at least	60 days prior to the date of	[movement]]		
		(2)	□ and/or	[II.2.2.1.2.	subjected to samples col		•		
		(2)	□ and/or	[II.2.2.1.3.	2.2.1.3. for at least 14 days prior to the date of movement and have b subjected to a PCR test, with negative results, carried out on s collected at least 14 days following the date of the commence the period of protection against attacks by vectors;]]]				
	(2)	□ and/or	[II.2.2.2.	2. have been kept at least for the 60 day period prior to departure in an establishment situated in a Member State or in an area of at least 150 km radio centred on the establishment, where surveillance in compliance with the requirements laid down in Sections 1 and 2 of Chapter 1 of Part II of Annex V to Delegated Regulation (EU) 2020/689 has been carried out during that period ar					
		(2)	□ either	[II.2.2.2.1.	24 of infects the past 2 y place where	ion with bluetongue virus v ears in an area of at least 1	inst those serotypes from 1 to which were reported during 50 km radius centred on the d are within the immunity s of the vaccine and		
			(2)	□ either	[II.2.2.2.1.1.	have been vaccinated more of movement]]]	re than 60 days before the date		
			(2)	□ and/or	[II.2.2.2.1.2.	subjected to a PCR test, wi	h an inactivated vaccine and ith negative results on samples after the onset of the immunity [the vaccine;]]]]		
		(2)	□ and/or	[II.2.2.2.2.	24 of infect the past 2 y	on with bluetongue virus	ninst those serotypes from 1 to which were reported during 50 km radius centred on the and		

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	II. Health info	ormation							
			(2)	o either	[II.2.2.2.2.1.	the animals have been subjet a serological test carried out 60 days before the date of m	on samples collected at least		
Certification			(2)	or	[II.2.2.2.2.2.	the animals have been subjet a serological test carried out 30 days before the date of the test, with negative results, callected not earlier than 14 movement.]]]]	on samples collected at least ne movement and to a PCR arried out on samples		
٠.	(2)(3) □ and/or	[II.2.2.	Part II of A	o not fulfil the requirements laid down in points 1 to 3 of Section 1 of Chapter 2 of Annex V to Regulation (EU) 2020/689 and the competent authority of the Mer f origin authorised movement of those animals to another Member State or zor					
(2) □ either [II.2.2.1. with the status free from infection with bluetongue virus (serot the Member State of destination has informed the Commission Member States that such movement is authorised subject to the referred to in Article 43(2)(a), (b) and (c) of Delegated Regulatio and									
		(2)	□ either	[II.2.2.1.1.		ection 1 of Chapter 2 of Part I Regulation, and	I of Annex V to that		
		(2)	□ and/or	[II.2.2.1.2.	-	ection 1 of Chapter 2 of Part I Regulation, and	I of Annex V to that		
		(2)	□ and/or	[II.2.2.1.3.	-	ection 1 of Chapter 2 of Part I Regulation, and	I of Annex V to that		
		(2)	□ and/or	[II.2.2.1.4.		ection 1 of Chapter 2 of Part I Regulation, and	I of Annex V to that		
					32(2) of Del	ments laid down in Article 32 egated Regulation (EU) 2020/0 n Article 33 of that Delegated	688 and the requirements		
	(2)	□ and/or	[II.2.2.2.	(serotypes Commission subject to	1-24) and the	ication program for infection e Member State of destination her Member States that such as referred to in Article 43(2)(89 and	n has informed the movement is authorised		
		(2)	\square either	[II.2.2.2.1.		ection 1 of Chapter 2 of Part I Regulation, and	I of Annex V to that		
		(2)	□ and/or	[II.2.2.2.2.	_	ection 1 of Chapter 2 of Part I Regulation, and	I of Annex V to that		
		(2)	□ and/or	[II.2.2.2.3.		ection 1 of Chapter 2 of Part I Regulation, and	I of Annex V to that		
		(2)	□ and/or	[II.2.2.2.4.	-	ection 1 of Chapter 2 of Part I Regulation, and	I of Annex V to that		
					32(2) of Del	ments laid down in Article 32 egated Regulation (EU) 2020/0 n Article 33 of that Delegated	688 and the requirements		
	(2)	□ and/or	[II.2.2.3.	by the erac 24) and the	dication prog e Member Sta	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		
		(2)	\square either	[II.2.2.3.1.	without any	y conditions, and			
		(2)	\square and/or	[II.2.2.3.2.		he conditions referred to in p of Annex V to Delegated Regu			
I			□ and/or			he conditions referred to in p			

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	II. Health info	rmation								
					2 of Part II	of Annex V to Delegated Re	gulation (E	U) 2020/689, and		
		(2)	□ and/or	[II.2.2.3.4.	,	ne conditions referred to in of Annex V to Delegated Re	•	-		
ے		(2)	□ and/or	[II.2.2.3.5.	•	ne conditions referred to in of Annex V to Delegated Re	-	_		
Part II: Certification					32(2) of Del	ments laid down in Article egated Regulation (EU) 202 n Article 33 of that Delegate	0/688 and t	the requirements		
ق ا	II.3	To the bes	t of my knov	wledge and a	as declared b	y the operator:				
Part II		II.3.1.				spatch there are no abnorr	nal mortali	ities with an		
		II.3.2.				act with animals which are .1., or with animals of a lov				
		II.3.3.	·							
	II.4.		ents are ma n (EU) 2020/6		ort the consi	gnment in accordance with	Article 4 o	of Delegated		
	II.5.	animals, t		te of issuing. In the case of of the certificate may be ex						
	lumpy skir		compliance			n relation to emergency pr d Annex IX, Part 3, point (3				
	skin diseas		iance with A			lation to emergency protec x IX, Part 3, point (3.2), to (
	Animal we	lfare attest	ation							
		ovisions of	*		,	th certificate were fit to be n the intended journey due				
	Notes:									
	In accordance with the Agreement on the withdrawal of the United Kingdom of Great Britain and Northern Irelan 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 certificate include the United Kingdom in respect of Northern Ireland.									
	This animal health certificate shall be completed according to the notes for the completion of certificates provided for in Chapter 2 of Annex I to Commission Implementing Regulation (EU) 2020/2235.									
	Part I:									
Box "Place of dispatch": Indicate a confined establishment a reference of Regulation (EU) 2016/429. I.11:						shment approved in accordance with Articles 97 and 99				
Box "Place of destination": Indicate a confined establishment approved in accordance will reference 99 of Regulation (EU) 2016/429. I.12:							ith Articles 97 and			
	Part II:									
	(1)	There can	be one or m	ore animals	s in the consi	gnment.				
	(2)	Delete if n	ot applicabl	e.						

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	II. Health information								
	(3) Only in case of animals belonging to the families Antilocapridae, Bovidae, Camelidae, Cervidae, Giraffidae, Moschidae or Tragulidae.								
	Certifying Officer/Official veterinarian Name (in capital letters)	Qualification and title							
tion	Date of signature Stamp	Signature							
Part II: Certification									
t II: Ce									
Par									

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