INTRA

	I.1. Consignor			I.2. IMSOC reference	I.2.a. Local reference	
	Name Address				I.3. Central Competent Authority	
	Country ISO Code				I.4. Local Competent Authority	
nt	I.5. Consignee			I.6. Operator conducting assembly operations independently of an establishment		
of consignment	Name			establishment Name		
gn	Address		700 O 1	Address		
nsi	Country		ISO Code	Approval		
CO]				Number Country	ISO Code	
				Country	130 code	
Part I: Description	I.7. Country of orig	I.7. Country of origin ISO Code I.8. Region of origin Code		I.9. Country of destination	ISO Code	
crip	I.8. Region of origi			I.10. Region of destination	Code	
es	I.11. Place of dispa	itch		I.12. Place of destination		
[: I	Name			Name		
rt .	Address			Address		
Pa	Approval Number			Approval Number		
	Country		ISO Code	Country	ISO Code	
	I.13. Place of loadi	nø		I.14. Date and time of departure		
	Name	116		1.14. Date and time of departure		
	Address					
	Approval					
	Nûmber Country ISO Code		ISO Code			
	, , , , , , , , , , , , , , , , , , ,			-		
	I.15. Means of Trai	International	Identification	I.16. Transporter Name		
	Mode	transport document	Tuentimeuton	Address		
		document		Approval Number		
				Number Country	ISO Code	
					130 code	
				I.17. Accompanying documents		
				Document Type Accompanying document		
				reference		
				Date of Issue		
				Country Place of issue		
	I.18. Transport cor	nditions		The or issue		
	Chilled □		Ambient \square	Frozen 🗆		
	I.19. Container Nu	mber / Seal Numbe	er			
	I.20. Certified as					
	Further processing	g 🗆	Further keeping \square	Germinal products \square	Dispatch centre \square	
	Registered equine animal \square		Travelling circus/animal act \square	Release into the wild \square	Live aquatic animals for human consumption \square	
	Organic fertilizers and soil improvers Other		Technical use	Exhibition \square	Event or activity near borders \Box	
			Products for human consumption	Ornamental aquaculture establishment	Relaying	
	Confined establish	mont [Quarantine or similar			
	Commed establish	инен 🗀	establishment	Slaughter		
	I.21. For transit through a third country					
	Third country		ISO Code			
	Exit point		BCP code			
	Entry point			BCP code		
	I.22. For transit through Member State(s)			I.23. For export		
	Member State ISO Code		ISO Code	Third country	ISO Code	
				Exit point	BCP code	
	I.24. Estimated journey time			I.25. Journey Log		

en 1/7

L27. Total quantity I.30. Description of consignment 1. 01 LIVE ANIMALS 0105 Live poultry, that is to say, fowls of the species Gallus domesticus, ducks, geese, turkeys and guinea fowls Weighing not more than 185 g: 010514 Geese 01051400 Geese #1. Commodity Species Age Quantity Identification system Identification Number Quantity						
1. 01 LIVE ANIMALS 0.105 Live noultry, that is to say, fowls of the species Gallus domesticus, ducks, goese, turkeys and guinea fowls						
0105 Live poultry, that is to say, fowls of the species Gallus domesticus, ducks, geese, turkeys and guinea fowls						
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Weighing not more than 185 g: 010514 Geese 01051400 Geese						
010514 Geese 01051400 Geese						
■ 01051400 Geese						
#1. Commodity Subcategory Identification system Identification Number	ar					
Species Age Quantity						
5 quantity						
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en 2/7

UNION 2024/1044 (2021/403) MODEL CONFINED-LIVE-INTR						
	II. Health information					
	II. Health i	nformation				
	I, the undersigned official veterinarian, hereby certify, that:					
	II.1.	The anima	ls (1) of the consignment described in Part I meet the following requirements:			
Part II: Certification	II.1.1.	Their confined establishment of dispatch is approved in accordance with Articles 97 and 99 of Regulation (EU) 2016/429 of the European Parliament and of the Council.				
	II.1.2.	They have not shown clinical signs or symptoms of diseases, in particular relevant diseases listed in the Annex to Commission Implementing Regulation (EU) 2018/1882, during the clinical examination, or where this is not possible, a clinical inspection, which was carried out within the last 48 hours prior to the time of departure of the consignment, on (insert date dd/mm/yyyy).				
	II.2.	_	to official information, animals of the consignment described in Part I meet the following uirements:			
Pe	II.2.1.	They come	from a confined establishment or a zone:			
	(2) o either	reasons of species, an	ct to movement restrictions affecting the species of animals to be moved and established for diseases listed for those species or diseases subject to emergency measures relevant for those d they have not been in contact with kept animals of a listed species of a lower health status quate period.]			
	(2) or	,	restrictions affecting the species of animals to be moved and established for cogations from movement restrictions have been granted, and:			
	(2)	☐ [they co	omply with the requirements set out in (4);]]			
	(2)	□ [and in	particular, they are (5).]]			
	(2) (6) □ either	[II.2.2.	They originate from a Member State or a zone thereof free from infection with bluetongue virus (serotypes 1-24), where no case of infection with bluetongue virus (serotypes 1-24) has been confirmed in the targeted animal population during the last 24 months prior to the date of departure of the consignment and have not been vaccinated with a live vaccine against infection with bluetongue virus (serotypes 1-24) in the last 60 days prior to the date of movement and the requirements laid down in Article 32(1), point (a), (b) or (c), or Article 32(2) of Commission Delegated Regulation (EU) 2020/688 are fulfilled.]			
	(2) (6) and/or	[II.2.2.	They originate from a Member State or a zone thereof covered by the eradication programme for infection with bluetongue virus (serotypes 1-24) and the requirements laid down in Article 32(1), point (a), (b) or (c), or Article 32(2) of Delegated Regulation (EU) 2020/688 are fulfilled, and they:			
	(2) 🗆 either	[II.2.2.1.	have been kept in a Member State or zone thereof seasonally free from infection with bluetongue virus (serotypes 1-24) in accordance with Article 40(3) of Commission Delegated Regulation (EU) 2020/689:			
	(2) □ either	[II.2.2.1.1.	for at least 60 days prior to the date of departure of the consignment;]]			
	(2) □ and/or	[II.2.2.1.2.	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 least 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);]]			
	(2) □ and/or	[II.2.2.1.3.	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.2.2.2.	have been protected against attacks by vectors during transportation to the place of destination and have been kept protected against attacks by vectors in a vector protected establishment:			
	(2) □ either	[II.2.2.2.1.	for at least 60 days prior to the date of departure of the consignment]]			
	(2) □ and/or	[II.2.2.2.2.	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 least 28 days following the date of the commencement of the period of protection against attacks			

en 3/7

-01	NION		2024/1044 (2021/403) MODEL CONFINED-LIVE-INTRA
	II. Health info	ormation	
			by vectors]]
u	(2) □ and/or	[II.2.2.2.3.	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 the commencement of the period of protection against attacks by vectors;]]]
Part II: Certification	(2) □ and/or	[II.2.2.3.	have been vaccinated against all serotypes from 1 to 24 of infection with bluetongue virus which were reported in that Member State or zone thereof during the last 2 years prior to the date of departure of the consignment and are within the immunity period guaranteed in the specifications of the vaccine, and:
t II: C	(2) □ either	[II.2.2.3.1.	have been vaccinated more than 60 days prior to the date of departure of the consignment;]]
Par	(2) □ and/or	[II.2.2.3.2.	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 date of onset of the immunity set in the specifications of the vaccine;]]]
	(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 in that Member State or zone thereof during the last 2 years prior to the date of departure of the consignment, and:
	(2) □ either	[II.2.2.4.1.	the serological test has been carried out on samples collected at least 60 days prior to the date of departure of the consignment;]]
	(2) □ and/or	[II.2.2.4.2.	the serological test has been carried out on samples collected at least 30 days prior to the date of departure of the consignment and the animals have been subjected to a PCR test, with negative results, carried out on samples collected not earlier than 14 days prior to the date of departure of the consignment;]]]
	(2) (6) and/or [II.2.2.	(serotypes (serotypes	nate from a Member State or a zone thereof neither free from infection with bluetongue virus 1-24) nor covered by the eradication programme for infection with bluetongue virus 1-24) and the requirements laid down in Article 32(1), point (a), (b) or (c), or Article 32(2) of Regulation (EU) 2020/688 are fulfilled, and they:
	(2) 🗆 either	[II.2.2.1.	have been protected against attacks by vectors during transportation to the place of destination and have been kept protected against attacks by vectors in a vector protected establishment:
	(2) □ either	[II.2.2.1.1.	for at least 60 days prior to the date of departure of the consignment;]]
	(2) □ and/or	[II.2.2.1.2.	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 least 28 days following the date of the commencement of the period of protection against attacks by vectors;]]
	(2) □ and/or	[II.2.2.1.3.	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 the commencement of the period of protection against attacks by vectors;]]]
	(2) □ and/or	[II.2.2.2.	have been kept at least for the last 60 days prior to the date of departure of the consignment in an establishment situated in a Member State or within an area of at least 150 km radius centred on the establishment, where surveillance in compliance with the requirements laid down in Part II, Chapter 1, Sections 1 and 2, of Annex V to Delegated Regulation (EU) 2020/689 has been carried out during that period, and:
	(2) □ either	[II.2.2.2.1.	the animals have been vaccinated 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 are within the immunity period guaranteed in the specifications of the vaccine, and:
	(2) □ either	[II.2.2.2.1. 1.	have been vaccinated more than 60 days prior to the date of departure of the consignment;]]]
	(2)		of the consignment in an area of at least 150 km radius centred on the animals were kept and are within the immunity period guaranteed in the vaccine, and: have been vaccinated more than 60 days prior to the date of departure

en 4/7

II. Health	information					
(2) □ and/or	[II.2.2.2.1. 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 date of onset of the immunity set in the specifications of the vaccine;]]]]				
(2) □ and/or	[II.2.2.2.2.	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.2.2.2.2. 1.	the animals have been subjected with positive results to a serological test carried out on samples collected at least 60 days prior to the date of departure of the consignment;]]]				
Part II: Certification (2) (2) (2) (2) (3) (4) (5) (6) (7) (7) (8) (9) (9) (1) (1) (1) (1) (1) (2) (3) (4) (5) (6) (7) (7) (8) (9) (9) (1) (1) (1) (1) (1) (2) (1) (2) (3) (4) (5) (6) (7) (7) (8) (8) (9) (9) (9) (9) (9) (9) (9) (9	r [II.2.2.2.2. 2.	the animals have been subjected with positive results to a serological test carried out on samples collected at least 30 days prior to the date of departure of the consignment and to a PCR test, with negative results, carried out on samples collected not earlier than 14 days prior to the date of departure of the consignment;]]]]				
	I.2. Delegated	ot fulfil the requirements laid down in Part II, Chapter 2, Section 1, points 1 to 3, of Annex V to Regulation (EU) 2020/689 and the competent authority of the Member State of origin movement of those animals to another Member State or zone thereof:				
(2) □ either	[II.2.2.1.	with the status "free from infection with bluetongue virus (serotypes 1-24)" and the Member State of destination has informed the Commission and the other Member States that such movement is authorised subject to the conditions referred to in Article 43(2), points (a), (b) and (c), of Delegated Regulation (EU) 2020/689 and:				
(2) □ either	[II.2.2.1.1.	Part II, Chapter 2, Section 1, point 5, of Annex V to that Delegated Regulation, and				
(2) □ and/or	[II.2.2.1.2.	Part II, Chapter 2, Section 1, point 6, of Annex V to that Delegated Regulation, and				
(2) □ and/or	[II.2.2.1.3.	Part II, Chapter 2, Section 1, point 7, of Annex V to that Delegated Regulation, and				
(2) □ and/or	[II.2.2.1.4.	Part II, Chapter 2, Section 1, point 8, of Annex V to that Delegated Regulation, and				
	the requirements laid down in Article 32(1), point (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	[II.2.2.2.	with an approved eradication program for infection with bluetongue virus (serotypes 1-24) and the Member State of destination has informed the Commission and the other Member States that such movement is authorised subject to the conditions referred to in Article 43(2), points (a), (b) and (c), of Delegated Regulation (EU) 2020/689, and:				
(2) □ either	[II.2.2.2.1.	Part II, Chapter 2, Section 1, point 5, of Annex V to that Delegated Regulation, and				
(2) □ and/or	[II.2.2.2.2.	Part II, Chapter 2, Section 1, point 6, of Annex V to that Delegated Regulation, and				
(2) □ and/or	[II.2.2.2.3.	Part II, Chapter 2, Section 1, point 7, of Annex V to that Delegated Regulation, and				
(2) □ and/or	[II.2.2.2.4.	Part II, Chapter 2, Section 1, point 8, of Annex V to that Delegated Regulation, and				
	the requirements laid down in Article 32(1), point (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	[II.2.2.3.	neither free from infection with bluetongue virus (serotypes 1-24) nor covered by the eradication programme for infection with bluetongue virus (serotypes 1-24) and the Member State of destination has informed the Commission and the other Member States that such movement is authorised:				
(2) □ either	[II.2.2.3.1.	without any conditions, and:				
(2) □ and/or	[II.2.2.3.2.	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				

en 5/7

U	NION		2024/1044 (2021/403) MODEL CONFINED-LIVE-INTRA	
	II. Health info	rmation		
	(2) □ and/or	[II.2.2.3.3.	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.2.2.3.4.	subject to the conditions referred to in Part II, Chapter 2, Section 1, point 7, of Annex V to Delegated Regulation (EU) 2020/689, and	
tion	(2) □ and/or	[II.2.2.3.5.	subject to the conditions referred to in Part II, Chapter 2, Section 1, point 8, of Annex V to Delegated Regulation (EU) 2020/689, and	
Cartification	the require 2020/688 a		down in Article 32(1), point (a), (b) or (c), or Article 32(2) of Delegated Regulation (EU) irements laid down in Article 33 of that Delegated Regulation are fulfilled.]]]	
2	II.3.	To the best	of my knowledge and as declared by the operator:	
ort III	II.3.1. II.3.2.		ined establishment of departure of the consignment there are no abnormal mortalities with mined cause affecting the animals to be moved.	
ם מ	II.3.2.		ls have not been in contact with animals which are subject to movement restrictions referred II.2.1, or with animals of a lower health status.	
	II.3.3.		he results of the surveillance plan of the confined establishment, the animals do not pose a risk at the confined establishment of destination for the spread of diseases for which they	
	II.4.		ents are made to transport the consignment in accordance with Article 4 of Delegated (EU) 2020/688.	
II.5. This animal health certificate is valid for 10 days from the date of issuing. In the case of transposition waterway/sea of animals, the period of 10 days for the validity of the certificate may be extended duration of the journey by waterway/sea.				
	Animal we	lfare attesta	ition	
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).				
	Notes:			
	In accordance with the Agreement on the withdrawal of the United Kingdom of Great Britain and Northern 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 animal health certificate include the United Kingdom in respect of Northern Ireland.			
	This animal health certificate shall be completed in accordance with the notes for the completion of certific 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 approved in accordance reference of Regulation (EU) 2016/429. I.11:		ispatch": Indicate a confined establishment approved in accordance with Articles 97 and 99 on (EU) 2016/429.		
Box "Place of destination": Indicate a confined establishment approver reference 99 of Regulation (EU) 2016/429. I.12:			estination": Indicate a confined establishment approved in accordance with Articles 97 and lation (EU) 2016/429.	
	Part II:			
	(1)	There may	be one or more animals in the consignment.	
	(2)	Delete if no	ot applicable.	
	(3)	Insert the 1	name of the disease(s).	
	(4)		specific reference to the article(s), titleand number of the relevant legal act(s) adopted by the on providing for those requirements.	
	(5)		specific attestation(s) provided for in and required by the relevant legal act(s) adopted by the on, as referred to in Article 126(1), points (b)(ii) and (iii), of Regulation (EU) 2016/429.	
(6) Only in the case of animals belonging to the families Antilocapridae, Bovidae, Camelidae, CGiraffidae, Moschidae or Tragulidae.				

en 6/7

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	II. Health information		
	Certifying Officer/Official veterinarian		
	Name (in capital letters)	Qualification and title	
	Date Stamp	Signature	
	Statitp		
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tio			
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tif			
Cer			
Part II: Certification			
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en 7/7