INTRA

	I.1. Consignor			I.2. IMSOC reference	I.2.a. Local reference		
	Name				I.3. Central Competent Authority		
	Address Country		ISO Code		I.4. Local Competent Authority		
of consignment	I.5. Consignee Name			I.6. Operator conducting assembly operations independently of an establishment			
틾	Address			Name			
Sig	Country		ISO Code	Address Approval			
				Number			
ij				Country	ISO Code		
	I.7. Country of orig	gin	ISO Code	I.9. Country of destination	ISO Code		
crip	I.8. Region of origi	in	Code	I.10. Region of destination	Code		
es	I.11. Place of dispa	ntch		I.12. Place of destination			
ᆲ	Name			Name			
티	Address			Address			
Pa	Approval Number			Approval Number			
	Country		ISO Code	Country	ISO Code		
ł	I.13. Place of loadi	ng		I.14. Date and time of departure			
	Name	11.6		1.14. Bute and time of departure			
	Address						
	Approval						
	Number Country		ISO Code				
- I	I.15. Means of Trai		I.J4:6:	I.16. Transporter			
	Mode	International transport	Identification	Name			
		document		Annroyal			
				Approval Number			
				Country	ISO Code		
				I.17. Accompanying documents			
				Document Type Accompanying document reference Date of Issue Country			
				Place of issue			
- 1	I.18. Transport cor	nditions					
	Ambient \square		Chilled \square	Frozen 🗆			
	I.19. Container Nu	mber / Seal Numbe	er				
	I.20. Certified as						
	Other 🗆		Further keeping \square	Exhibition \square	Confined establishment \square		
	Release into the w	rild 🗆	Travelling circus/animal act \square	Breeding \square	Quarantine or similar establishment \square		
	Event or activity near borders						
f	I.21. For transit through a third country						
	Third country			ISO Code			
	Exit point Entry point			BCP code BCP code			
ŀ	I.22. For transit th	rough Mombor Cto	to(c)	I.23. For export			
		rough member sta	_				
	Member State		ISO Code	Third country Exit point	ISO Code BCP code		
	I.24. Estimated jou	ırney time		I.25. Journey Log			
- 1	I.27. Total quantity			I.28. Total gross weight			
ŀ	I.30. Description of			1.20. 2011 51000 1101511			
	_	-					
ļ	1. 01 LIVE ANIMA	LO					

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	0104 Live sheep and goats						
	010410 Sheep #1. Commodity	Cubactagamy	Corr	Identification avatem			
			Sex	Identification system			
	Species	Identification Number	Age	Quantity			
int							
m							
gn							
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I: I							
Part I: Description of consignment							
Pē							

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_	UNIUN				2024/1044 (202	1/403)	MODEL OV/CAI	P-INTRA-X
Part II: Certification	II. Health in	formation						
	II. Health	informatio	n					
		I, the undersigned official veterinarian, hereby certify that:						
	II.1.	II.1. The ovine/caprine animals (1) of the consignment described in Part I meet the following requirements:						
	II.1.1.		identified as on (EU) 2019/	provided for in Article 4 2035.	5(2) or (4) or Article 4	6(1) of (Commission Deleg	gated
	II.1.2.	-	at least 30 da than 30 days	ays prior to the date of do of age,	eparture of the consig	nment,	or since birth, if t	hey are
	II.1.2.1.	have bee	n continuous	sly resident in the establi	shment of origin;			
	II.1.2.2.	II.1.2.2. have not been in contact with kept ovine or caprine animals of a lower health status or subject to movement restrictions for animal health reasons;						
	II.1.2.3.	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 consignment.						
	II.1.3.							
	II.2.	Accordin requirem	_	nformation, the animals	described in Part I me	eet the f	ollowing health	
L	— II.2.1.	(2)	o either	[They come from estab affecting ovine/caprine those species or diseas species, and they have of a lower health status	e animals and establis es subject to emergen not been in contact w	hed for cy meas ith kept	reasons of diseas ures relevant for	es listed for those
	(2)	o or	ovine/cap	ne from establishments o rine animals and establis t restrictions have been a	shed for		restrictions affect derogations fron	-
	(2)	□ [they	comply with	the requirements set out	in (4);]]		
	(2)	□ [and i	n particular,	they are (5	5).]]			
	(2) o either	[II.2.2.	-	e from establishments fr hout vaccination regard				itensis and
	(2) □ either			f origin are situated in a a a abortus, B. melitensis a				
(2) [they have been subjected to a test for infection with Brucella abort and/or of the diagnostic methods provided for in Part 1 of Annex I to Comm 2020/688, carried out, with negative results, on a sample taken durit of departure of the consignment, and in the case of post-parturient parturition;]]				nission I ng the la	Delegated Regulat ast 30 days prior t	ion (EU) o the date		
	(2) □ and/or	[they are	less than 6 n	nonths old;]]				
	(2) □ and/or							
	(2) o or	[II.2.2.	B. suis wit Member S	e from establishments fr h vaccination regarding tate or zone thereof with sis and B. suis regarding	ovine and caprine and nout the status free fro	imals ar om infec	nd they are moved	d to a
	(2) □ either	[II.2.3.	Mycobacto	kept ovine animals and c erium tuberculosis comp luring the last 42 days pr	lex (M. bovis, M. capr	ae and l	M. tuberculosis) h	
	(2) □ and/or	[II.2.3.	infection v tuberculos at least 12	kept caprine animals and with Mycobacterium tub sis) has been carried out months prior to the date elegated Regulation (EU)	erculosis complex (M. on the caprine anima e of departure of the c	bovis, l ls kept i	M. caprae and M. In the establishme	ents during

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O1	NIOIN		2024/1044 (2021/403)	MODLL OV/CAP-INTRA-A				
	II. Health info	ormation						
	II.2.4.	They come from establishments in which infection been reported during the last 30 days prior to	_					
	II.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:						
: Certification	(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 the date of departure of the consignment and the animals have been kept in a zone seasonally free from epizootic haemorrhagic disease in accordance with Parts 1 and 2 of Annex IX to Delegated Regulation (EU) 2020/688;						
Part II:	(2) □ either	for at least 60 days prior to the date of departure of the consignment;]						
	(2) □ and/or	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 seasonally disease-free area;]						
	(2) □ and/or	for at least 14 days prior to the date of departutest, with negative results, carried out on samp the animals into the seasonally disease-free ar	oles collected at least 14 days	•				
	(2) □ and/or	[has been reported in kept animals of listed sp date of departure of the consignment and the during transportation to the place of destination vectors in a vector protected establishment full IX to Delegated Regulation (EU) 2020/688;	animals have been protected on and they have been kept p	against attacks by vectors rotected against attacks by				
	(2) □ either	for at least 60 days prior to the date of departu	re of the consignment;]					
	(2) □ and/or	for at least 28 days prior to the date of departuserological test, with negative results, carried date of the commencement of the period of prior to the date of departusers.	out on samples collected at le	ast 28 days following the				
	(2) □ and/or	for at least 14 days prior to the date of departutest, with negative results, carried out on samp commencement of the period of protection again	oles collected at least 14 days					
	(2) 🗆 and/or	[has been reported in kept animals of listed sp date of departure of the consignment and the Commission and the other Member States that	Member State of destination	has informed the				
	II.2.6.	They come from establishments in which anth days prior to the date of departure of the const	_	n reported during the last 15				
	II.2.7.	They come from establishments in which surr last 30 days prior to the date of departure of the		not been reported during the				
	(2) o either	[surra has not been reported in the establishm of the consignment.]	nents during the last 2 years p	orior to the date of departure				
(2) ∘ or		[surra has been reported during the last 2 year following the date of the last outbreak, the afferestrictions until the date on which the infecte and the remaining animals in the establishmediagnostic methods provided for in Part 3 of A with negative results, on samples taken at least animals have been removed from the establishmediagnostic methods animals have been removed from the stablishmediagnostic methods.	ected establishments have reset animals have been remove nts have been subjected to a tennex I to Delegated Regulations of months following the date	mained under movement d from the establishments, test for surra with one of the on (EU) 2020/688, carried out,				
	(2)	\square [II.2.8. They are kept uncastrated male ov	rine animals, and:					
	_	come from establishments in which ovine epic last 12 months prior to the date of departure o		not been reported during the				
	_	have been subjected to a serological test for ownegative results, on a sample taken during the consignment.]						

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	NION		2024/1044 (2021/403) MODEL 07/CAF-INTRA-2			
	II. Health info	ormation				
ation	(2) □ either	[II.2.9.	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 late of departure of the consignment and have not been vaccinated with a live vaccine against infection with bluetongue virus (serotypes 1-24) within the last 60 days prior to the late of departure of the consignment and the requirements laid down in Article 32(1), points (a), (b) or (c), or Article 32(2) of Delegated Regulation (EU) 2020/688 are fulfilled.]			
Part II: Certification	(2) 🗆 and/or	[II.2.9.	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), points (a), (b) or (c), or Article 32(2) of Delegated Regulation (EU) 2020/688 are fulfilled, and they:			
Part	(2) □ either	[II.2.9.1.	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.2.9.1.1.	for at least 60 days prior to the date of departure of the consignment;]]			
	(2) 🗆 and/or	[II.2.9.1.2.	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 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);]]			
	(2) 🗆 and/or	[II.2.9.1.3.	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.2.9.2.	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.2.9.2.1.	for at least 60 days prior to the date of departure of the consignment;]]			
	(2) 🗆 and/or	[II.2.9.2.2.	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 leas 28 days following the date of the commencement of the period of protection against attacks by vectors;]]			
	(2) 🗆 and/or	[II.2.9.2.3.	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;]]]			
	(2) \square and/or	[II.2.9.3.	have been vaccinated against all serotypes from 1 to 24 of infection with bluetongue virus which were reported during the last 2 years in that Member State or zone thereof and are within the immunity period guaranteed in the specifications of the vaccine, and:			
	(2) □ either	[II.2.9.3.1.	have been vaccinated more than 60 days prior to the date of departure of the consignment;]			
	(2) □ and/or	[II.2.9.3.2.	have been vaccinated with an inactivated vaccine and subjected to a PCR test, with negative results, carried out on samples collected at least 14 days after the date of the onset of the immunity set in the specifications of the vaccine;]]]			
	(2) 🗆 and/or	[II.2.9.4.	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 last 2 years prior to the date of departure of the consignment in that Member State or zone thereof, and:			
	(2) □ either	[II.2.9.4.1.	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.2.9.4.2.	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			

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	NIOIN			2024/1044 (2021/403)	•	
	II. Health in	formation				
			date of departure of the consignment			
u	(2) □ and/or	[II.2.9.	They originate from a Member State bluetongue virus (serotypes 1-24) no with bluetongue virus (serotypes 1-point (a), (b) or (c), or Article 32(2) or	e or a zone thereof neither for or covered by the eradication 24) and the requirements lai	n programme for infection d down in Article 32(1),	
Certification	(2) 🗆 either	[II.2.9.1.	have been protected against attacks destination and have been kept pro establishment:			
	(2) □ either	[II.2.9.1.1.	for at least 60 days prior to the date	of departure of the consign	ment;]]	
Part II:	(2) □ and/or	[II.2.9.1.2.	subjected to a serological test, with	for at least 28 days prior to the date of departure of the consignmentand 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.2.9.1.3.	for at least 14 days prior to the date of departure of the consignmentand 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.2.9.2.	have been kept for the last 60 days pestablishment situated in a Member the establishment, where surveillar II, Chapter 1, Sections 1 and 2, of Ancarried out during that period, and:	r State or in an area of at lea nce in compliance with the ro nnex V to Delegated Regulation	st 150 km radius centred on equirements set out in Part	
	(2) □ either	[II.2.9.2.1.	the animals have been vaccinated a bluetongue virus which were repor of the consignment in an area of at animals were kept and are within the the vaccine and:	ted during the last 2 years p least 150 km radius centred	rior to the date of departure on the place where the	
	(2) □ either	[II.2.9.2.1. 1.	have been vaccinated more than 60 consignment;]]]	days prior to the date of dep	parture of the	
	(2) 🗆 and/or	[II.2.9.2.1. 2.	have been vaccinated with an inact results, carried out on samples colle immunity set in the specifications o	ected at least 14 days after th		
	(2) 🗆 and/or	[II.2.9.2.2.	the animals have been immunised a bluetongue virus which were repor of the consignment in an area of at animals were kept, and:	ted during the last 2 years p	rior to the date of departure	
	(2) \square either	[II.2.9.2.2. 1.	the animals have been subjected wi samples collected at least 60 days pr	-	O .	
	(2) 🗆 and/or	[II.2.9.2.2. 2.	the animals have been subjected wi samples collected at least 30 days pr PCR test, with negative results, carr, prior to the date of departure of the	rior to the date of departure ied out on samples collected	of the consignment and to a	
	(2) 🗆 and/or	[II.2.9.	They do not fulfil the requirements Annex V to Delegated Regulation (E State of origin authorised movement thereof:	U) 2020/689 and the compete	ent authority of the Member	
	(2) □ either	[II.2.9.1.	with the status free from infection v State of destination has informed th movement is authorised subject to t and (c), of Delegated Regulation (EU	ne Commission and the other the conditions referred to in	Member States that such	
	(2) 🗆 either	[II.2.9.1.1.	Part II, Chapter 2, Section 1, point 5,	, of Annex V to that Delegate	d Regulation, and	

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	II. Health	information					
Certification	(2) □ and/or	[II.2.9.1.2.	Part II, Chapter 2, Section 1, point 6	6, of Annex V to that Delegated	d Regulation, and		
	(2) \square and/or	[II.2.9.1.3.	Part II, Chapter 2, Section 1, point 7	, of Annex V to that Delegated	d Regulation, and		
	(2) and/or	[II.2.9.1.4.	Part II, Chapter 2, Section 1, point 8	8, of Annex V to that Delegated	d Regulation, and		
	the requ 2020/68	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) \square and/or	[II.2.9.2.	24) and the Member State of destin Member States that such movemen	with an approved 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 under the conditions referred to in Article 43(2), points (a), (b) and (c), of Delegated Regulation (EU) 2020/689, and:			
	(2) □ either	[II.2.9.2.1.	Part II, Chapter 2, Section 1, point 5, of Annex V to that Delegated Regulation, and				
	(2) \square and/or	[II.2.9.2.2.	Part II, Chapter 2, Section 1, point 6	Part II, Chapter 2, Section 1, point 6, of Annex V to that Delegated Regulation, and			
	(2) □ and/or	[II.2.9.2.3.	Part II, Chapter 2, Section 1, point 7	, of Annex V to that Delegated	d Regulation, and		
L	(2) \square and/or	[II.2.9.2.4.	Part II, Chapter 2, Section 1, point 8	8, of Annex V to that Delegated	d Regulation, and		
			down in Article 32(1), points (a), (b) irements laid down in Article 33 of t				
	(2) □ and/or	[II.2.9.3.	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.2.9.3.1.	without any conditions, and				
	(2) \square and/or	[II.2.9.3.2.	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) \square and/or	[II.2.9.3.3.	. under the conditions referred to in Part II, Chapter 2, Section 1, point 6, of Annex V to Delegated Regulation (EU) 2020/689, and				
	(2) \square and/or	[II.2.9.3.4.	[II.2.9.3.4. under the conditions referred to in Part II, Chapter 2, Section 1, point 7, of Annex V to Delegated Regulation (EU) 2020/689, and				
	(2) \square and/or	[II.2.9.3.5.	under the conditions referred to in Delegated Regulation (EU) 2020/689	-	point 8, of Annex V to		
	_		down in Article 32(1), points (a), (b) irements laid down in Article 33 of t		_		
	(2) o either	[II.2.10.	The animals are intended for a Meropoint 2.3, of Annex VIII to Regulation the Council as having a negligible relisted in Chapter A, Section A, point having an approved national scrap	on (EC) No 999/2001 of the Eurisk status for classical scrapid t 3.2, of Annex VIII to Regulat	ropean Parliament and of e or for a Member State		
(2) [come from a holding situated in a Member State or zone the either 2.3, of Annex VIII to Regulation (EC) No 999/2001 as having							
(2) [come from a holding recognised as having a negligible ri and/or Chapter A, Section A, point 1.2, of Annex VIII to Regulation competent authority of the Member State in accordance v VIII to Regulation (EC) No 999/2001.]				Regulation (EC) No 999/2001	and listed as such by the		
(2) [come from a holding not subject to the measures laid down in Chapter B, points 3 and/or Regulation (EC) No 999/2001 and the animals are of the ovine species and are of the protein genotype, or the animals are of the caprine species and carry at least one of the caprine species.					re of the ARR/ARR prion		

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	NION			2024/1044 (2021/403)	WIGDEE OVICIN INVITATION		
	II. Health information						
		S146 alleles.]		I		
	(2) □ and/or		and are destined for a confined est EU) 2016/429 of the European Parl:	stablishment as defined in Article 4, point (48), of liament and of the Council.]			
Part II: Certification	(2) □ or	[comply with No 999/2001	h the conditions set out in Chapter .]]	A, Section A, point 4.1(d), of A	Annex VIII to Regulation (EC)		
	(2) ∘ or	t 9	[II.2.10. The animals are for breeding and are intended for a Member State or zone thereof other than those listed in Chapter A, Section A, point 2.3, of Annex VIII to Regulation (EC) No 999/2001 as having a negligible risk status for classical scrapie or other than those listed in Chapter A, Section A, point 3.2, of Annex VIII to Regulation (EC) No 999/2001 as having an approved national scrapie control programme, and:				
Part I	(2) 🗆 either	[come from a holding situated in a Member State or zone of a Member State listed in Chapter A, Section A, point 2.3, of Annex VIII to Regulation (EC) No 999/2001 as having a negligible risk status for classical scrapie.]					
	(2) □ and/or	[come from a holding recognised as having a negligible risk of classical scrapie in accordance with Chapter A, Section A, point 1.2, of Annex VIII to Regulation (EC) No 999/2001 and listed as such by the competent authority of the Member State in accordance with Chapter A, Section A, point 1.1, of Annex VIII to Regulation (EC) No 999/2001.]					
	(2) □ and/or	Chapter A, S competent a	a holding recognised as having a c ection A, point 1.3, of Annex VIII to outhority of the Member State in ac lation (EC) No 999/2001.]	Regulation (EC) No 999/2001	and listed as such by the		
	(2) □ and/or	Regulation (a holding not subject to the measu EC) No 999/2001 and the animals a otype, or the animals are of the cap]	re of the ovine species and a	re of the ARR/ARR prion		
	(2) \square [come from and are destined for a confined and/or Regulation (EU) 2016/429.]			stablishment as defined in Article 4, point (48), of			
	(2) □ or	(2) \square or [comply with the conditions set out in Chapte No 999/2001.]]		r A, Section A, point 4.1(d), of Annex VIII to Regulation (EC)			
	than those listed in Chapter A, Se 999/2001 as having a negligible r Chapter A, Section A, point 3.2, o		The animals are not for breeding a than those listed in Chapter A, Sect 1999/2001 as having a negligible risl Chapter A, Section A, point 3.2, of A approved national scrapie control	ion A, point 2.3, of Annex VII c status for classical scrapie o unnex VIII to Regulation (EC)	I to Regulation (EC) No r other than those listed in		
	II.3.		of my knowledge and as declared be were no abnormal mortalities wit	by the operator, the animals come from establishments ith an undetermined cause.			
	II.4.		nts are made to transport the consi EU) 2020/688.	gnment in accordance with A	article 4 of Delegated		
	 II.5. This animal health certificate is valid for 10 downtown waterway/sea of animals, the period of validity journey by waterway/sea. (2) (6) Since the date of departure from their establishment approved for assembly operation undergone more than two assembly operations. (2) (2) (2) (2) (2) (2) (3) (4) (4) (4) (4) (4) (4) (4) (4) (4) (4		ea of animals, the period of validity	-			
			ns, none of the animals of the				
			from their establishments of origin	.]]			
	(2) o or	[at least one establishme	of the animals of the consignment nt.]]	has undergone one assembly	y operation in an approved		
	(2) ∘ or	[at least one establishme	of the animals of the consignment nts.]]	has undergone two assembl	y operations in the approved		
	Animal we	lfare attestati	ion				
	At the time	At the time of inspection, the animals covered by this animal health certificate were fit to be transported in					

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UNION 2024/1044 (2021/403) MODEL OV/CAP-INTRA-X II. Health information accordance with the provisions of Council Regulation (EC) No 1/2005 on the intended journey due to start on (insert date) (7) (8). 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 the Union in this 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 certificates provided for in Chapter 2 of Annex I to Commission Implementing Regulation (EU) 2020/2235. Part I: Box "Place of dispatch": Indicate an establishment of the origin of the animals in the consignment or an reference establishment approved for assembly operations in accordance with Articles 97 and 99 of Regulation (EU) 2016/429. I.11: "Place of destination": Indicate an establishment of the final destination of the consignment or an Box establishment approved for assembly operations in accordance with Articles 97 and 99 of Regulation reference I.12: (EU) 2016/429. 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, shall be indicated. Box "Identification number": Indicate identification codes of the animals in the consignment identified in reference accordance with Article 45(2) or (4) or Article 46(1) of Delegated Regulation (EU) 2019/2035. I.30: Part II: (1) There may be one or more animals in the consignment. (2) Delete if not applicable. (3) Insert the name of the disease(s). (4)Insert the specific reference to the article(s), title, and number of the relevant legal act(s) adopted by the Commission providing for those requirements. Insert the specific attestation(s) provided for in and required by the relevant legal act(s) adopted by the (5) Commission, as referred to in Article 126(1), points (b)(ii) and (iii), of Regulation (EU) 2016/429. (6)Applicable in case the consignment is dispatched from the establishment approved for assembly operations. In the case where a consignment is grouped in an establishment approved for assembly operations and (7)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. (8) This statement does not exempt transporters from their obligation in accordance with Union rules in force in particular regarding the fitness to be transported. Certifying Officer/Official veterinarian Name (in capital letters) Oualification and title Signature Date Stamp

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