**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	
of consignment	I.5. Consignee			I.6. Operator conducting assembly operations independently of an establishment		
	Name Address			Name		
sig	Country		ISO Code	Address		
ons				Approval Number		
$^{\mathrm{jc}}$				Country	ISO Code	
	I.7. Country of orig	gin	ISO Code	I.9. Country of destination	ISO Code	
rip	I.8. Region of origi	in	Code	I.10. Region of destination	Code	
)es	I.11. Place of dispa	atch		I.12. Place of destination		
$\mathbf{I}$ :	Name			Name		
ır	Annexal			Address		
Pe	Approval Number			Approval Number		
	Country		ISO Code	Country	ISO Code	
	I.13. Place of loadi	ng		I.14. Date and time of departure		
	Name					
_	Address Approval					
	Number		700 0 1			
	Country		ISO Code			
	I.15. Means of Tra		T-1	I.16. Transporter		
	Mode	International transport	Identification	Name Address		
		document		_		
				Approval Number	100 0-1-	
				Country	ISO Code	
				I.17. Accompanying documents		
				Document Type Accompanying document		
				reference		
				Date of Issue		
				Country Place of issue		
	I.18. Transport con	nditions				
	Chilled $\square$		Ambient $\square$	Frozen $\square$		
	I.19. Container No	/ Seal No				
	I.20. Certified as					
	Travelling circus/a	animal act $\square$	Exhibition	Release into the wild $\square$	Slaughter	
	Other $\square$		Further keeping $\square$	Confined establishment $\square$	Event or activity near borders $\Box$	
	Quarantine or sime establishment $\square$	ilar				
	I 21 For transit th	rough a third coun	try			
	Third country	rough a thiru coun	шу	ISO Code		
	Exit point			BCP code		
	Entry point			BCP code		
	I.22. For transit th	rough Member Sta	te(s)	I.23. For export		
	Member State		ISO Code	Third country	ISO Code	
				Exit point	BCP code	
	194 Pastar 1 *	arr ti		I.25. Journey Log		
	I.24. Estimated jou					
	I.27. Total quantity	у		I.28. Total gross weight		
		y f consignment				

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	NION			INTRA
	0106 Other live animals			
	Mammals:			
	<b>010619</b> Other			
	<b>01061900</b> Other			
	#1. Commodity	Sex	Identification system	Identification Number
Part I: Description of consignment	Charies	Quantity		
<u>a</u>	Species	Quantity	Age	
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	II. Health inform	ation								
	I, the undersi	igned officia	l veterinari	an, hereby certify tha	t:					
	II.1.			consignment describe ne, camelid and cervic	_	_	other than bovine, ovine, ing requirements:			
	II.1.1.	They are identified as provided for in Article 117 of Regulation (EU) 2016/429 of the European Parliament and of the Council.								
Part II: Certification	II.1.2.	They, for at younger th			departure of the co	onsignmen	t, or since birth, if they are			
rti	II.1.2.1.	have been	continuous	ly resident in the estal	olishment of origin;					
: II: Ce	II.1.2.2.		have not been in contact with other kept ungulates of a lower health status or subject to movement restrictions for animal health reasons;							
Part	II.1.2.3.			t or indirect contact w ıring the last 30 days ı	_		tered the Union from a third of the consignment.			
	II.1.3.	They have not shown clinical signs or symptoms of diseases listed for ungulates of the species concerned during the clinical examination which was carried out, within 24 hours prior to the time of departure of the consignment, on (insert date dd/mm/yyyy).								
	II.2.	According to		nformation, the anima	ls described in Part	I meet the	e following health			
	II.2.1.	(2)	o either	reasons of diseases l measures relevant fo	the species of aning isted for those specior or those species, and	nals to be i ies or dise d they hav	ject to movement moved and established for ases subject to emergency e not been in contact with tatus for an adequate			
	(2)	$\circ$ or	species of	ne from establishment animals to be moved ement restrictions hav	and established for		nt restrictions affecting the (3), but derogations			
	(2)	☐ [they comply with the requirements set out in (4);]]								
	(2)	☐ [and in ]	particular. t	they are	(5).]]					
	II.2.2.			lishments in which an of departure of the co		has not be	en reported during 15 days			
	(2)	□ [II.2.3.	and B. sui	some from establishments in which infection with Brucella abortus, B. melitensis suis in kept animals of listed species has not been reported during the last 42 days to the date of departure of the consignment.]						
	(2)	□ [II.2.4.	complex (	They come from establishments in which infection with Mycobacterium tuberculosis complex (M. bovis, M. caprae and M. tuberculosis) in kept animals of listed species has not been reported during the last 42 days prior to the date of departure of the consignment.]						
	(2)	□ [II.2.5.		as not been reported o			es virus in kept terrestrial o the date of departure of the			
	(2)	□ [II.2.6.	-				least 150 km radius around orrhagic disease virus:			
	(2) □ either			l in kept animals of lis of the consignment;]	ted species for that	disease di	uring the last 2 years prior to			
	(2) □ and/or	date of dep epizootic h	arture of th aemorrhag		e animals have bee	n kept in a	g the last 2 years prior to the a zone seasonally free from x IX to Commission			
		(2) □ either	for at leas	t 60 days prior to the o	late of departure of	the consi	gnment;]			
		(2) □ and/or					gnment and have been ut on samples collected at			

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	ION		2021/1011	(2021/403) Model O1111	- CINGULATED INTIGERAL
	II. Health inform	ation			
			least 28 days following the date area;]	of entry of the animals into t	he seasonally disease-free
n		(2) □ and/or	for at least 14 days prior to the d subjected to a PCR test, with neg days following the date of entry	ative results, carried out on	samples collected at least 14
=	(2) □ and/or	date of dep during tran vectors in a	eported in kept animals of listed sarture of the consignment and the sportation to the place of destinance vector protected establishment fated Regulation (EU) 2020/688;	e animals have been protect tion and they have been kep	ed against attacks by vectors t protected against attacks by
art II		(2) □ either	for at least 60 days prior to the d	late of departure of the cons	ignment;]
P		(2) □ and/or	for at least 28 days prior to the d subjected to a serological test, w least 28 days following the date attacks by vectors;]	ith negative results, carried	out on samples collected at
		(2) □ and/or	for at least 14 days prior to the d subjected to a PCR test, with neg days following the date of comm vectors;]	ative results, carried out on	samples collected at least 14
	(2) □ and/or	date of dep	eported in kept animals of listed s arture of the consignment and th n and the other Member States th	e Member State of destination	n has informed the
	(2)	□ [II.2.7.	They come from establishments reported during the last 30 days		
	(2) o either		not been reported in the establish of the consignment.]]	ments during the last 2 year	s prior to the date of
	(2) ∘ or	the date of until the da remaining diagnostic i out, with no	the last 2 year the last outbreak, the affected est the on which the infected animals animals in the establishments have methods provided for in Part 3 of egative results, on samples taken we been removed from the establish	ablishments have remained have been removed from th ve been subjected to a test fo Annex I to Delegated Regula at least 6 months following t	under movement restrictions e establishments, and the r surra with one of the tion (EU) 2020/688, carried
	(2) □ either	[II.2.8.	They originate from a Member S bluetongue virus (serotypes 1-24) has been confirmenths prior to the date of departmenth a live vaccine against infect 60 days prior to the date of departmenth and	<ol> <li>where no case of infection med in the targeted animal parture of the consignment an tion with bluetongue virus (surture of the consignment an</li> </ol>	a with bluetongue virus population during the last 24 d have not been vaccinated serotypes 1-24) during the last d the requirements laid
	(2) □ and/or	[II.2.8.	They originate from a Member S programme for infection with blaid down in Article 32(1), point 2020/688 are fulfilled, and:	luetongue virus (serotypes 1	-24) and the requirements
		(2) □ either	infection with blueto	Member State or zone thered Ingue virus (serotypes 1-24) i Delegated Regulation (EU) 2	n accordance with Article
	(2) □ either	[II.2.8.1.1.	for at least 60 days prior to the d	late of departure of the cons	ignment;]]
	(2) □ and/or	[II.2.8.1.2.	for at least 28 days prior to the d subjected to a serological test, w least 28 days following the date thereof seasonally free from info	ith negative results, carried of entry of the animals into t	out on samples collected at he Member State or zone

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_	INION		2024/1044 (2021/403) Model Office-UNGULATES-INTRA-X
	II. Health inform	ation	
	(2) □ and/or	[II.2.8.1.3.	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);]]]
•	cauon	(2) □ and/or	[II.2.8.2. 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:
١	(2) □ either	[II.2.8.2.1.	for at least 60 days prior to the date of departure of the consignment;]]
1	(2) □ either (2) □ and/or	[II.2.8.2.2.	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 by vectors;]]
	(2) □ and/or	[II.2.8.2.3.	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) □ and/or	[II.2.8.3. 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:
	(2) □ either	[II.2.8.3.1.	have been vaccinated more than 60 days prior to the date of departure of the consignment;]]
	(2) 🗆 and/or	[II.2.8.3.2.	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.2.8.4. 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.2.8.4.1.	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.2.8.4.2.	the serological test has been carried out on samples collected at least 30 days prior to the date of the 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) □ and/or	[II.2.8.	They originate from a Member State or a zone thereof 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 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.2.8.1. 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.2.8.1.1.	for at least 60 days prior to the date of departure of the consignment;]]
	(2) □ and/or	[II.2.8.1.2.	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.2.8.1.3.	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 during at least 14 days following the date of the commencement of the period of protection against attacks by vectors;]]]
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	II. Health inform	ation					
80		(2) □ and/or	[II.2.8.2.	of at least 150 km rad compliance with the	tablishment situated i lius centred on the est requirements set out i gated Regulation (EU)	n a Me ablishr n Part	te of departure of the mber State or within an area nent, where surveillance in II, Chapter 1, Sections 1 and 89 has been carried out
Dawt II. Contification	(2) □ either	[II.2.8.2.1.	bluetongue departure where the	e virus which were rep of the consignment in	oorted during the last an area of at least 150 d are within the immu	2 years ) km ra	1 to 24 of infection with s prior to the date of dius centred on the place eriod guaranteed in the
Daret	(2) □ either	[II.2.8.2.1.1.	have been consignme	vaccinated more than nt;]]]	60 days prior to the d	late of o	departure of the
	(2) □ and/or	[II.2.8.2.1.2.	negative re	vaccinated with an in esults, carried out on s e immunity set in the	amples collected at le	ast 14 d	lays after the date of the
	(2) □ and/or	[II.2.8.2.2.	bluetongue departure	e virus which were rep	ported during the last thin an area of at leas	2 years	1 to 24 of infection with s prior to the date of m radius centred on the
	(2) □ either	[II.2.8.2.2.1.			_		ological test carried out on re of the consignment;]]]
	(2) □ and/or	[II.2.8.2.2.2.	samples co with negat	llected at least 30 days	s prior to the date of t t on samples collected	he mov	rological test carried out on rement and to a PCR test, rlier than 14 days prior to
	(2) □ and/or[II.2.8.	to Delegated	d Regulation		e competent authority	y of the	n 1, points 1 to 3, of Annex V Member State of origin thereof:
		(2) □ either	[II.2.8.1.	the Member State of o	destination has inforn uch movement is auth	ned the norised	te virus (serotypes 1-24) and Commission and the other subject to the conditions of Delegated Regulation (EU)
	(2) □ either	[II.2.8.1.1.	Part II, Cha	pter 2, Section 1, poin	t 5, of Annex V to that	Regula	ation, and
	(2) □ and/or	[II.2.8.1.2.	Part II, Cha	pter 2, Section 1, poin	t 6, of Annex V to that	Regula	ation, and
	(2) □ and/or	[II.2.8.1.3.	Part II, Cha	pter 2, Section 1, poin	t 7, of Annex V to that	Regula	ation, and
	(2) □ and/or	[II.2.8.1.4.	Part II, Cha	pter 2, Section 1, poin	t 8, of Annex V to that	Regula	ation, and
				e 32(1), point (a), (b) o lown in Article 33 of tl			
		☐ (2) and/or	[II.2.8.2.	(serotypes 1-24) and t	the Member State of do ther Member States to ons referred to in Arti	estinat that suc	ection with bluetongue virus ion has informed the ch movement is authorised 2), points (a), (b) and (c), of
	(2) □ either	[II.2.8.2.1.	Part II, Cha	pter 2, Section 1, poin	t 5, of Annex V to that	Delega	ated Regulation, and
	(2) □ and/or	[II.2.8.2.2.	Part II, Cha	apter 2, Section 1, poin	t 6, of Annex V to that	Delega	ated Regulation, and
	(2) □ and/or	[II.2.8.2.3.	Part II, Cha	apter 2, Section 1, poin	t 7, of Annex V to that	Delega	ated Regulation, and

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O1	11011			2024/104	4 (2021/403) MOU	ci Ollic	R-UNGULATES-INTRA-A		
	II. Health inform	ation							
	(2) □ and/or	[II.2.8.2.4.	Part II, Cha	apter 2, Section 1, po	int 8, of Annex V to	that Delega	ated 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;]]]									
Part II: Certification		(2) □ and/or	[II.2.8.3.	by the eradication j 1-24) and the Meml	orogramme for infe	ction with ion has inf	(serotypes 1-24) nor covered bluetongue virus (serotypes ormed the Commission and uthorised:		
er	(2) □ either	[II.2.8.3.1.	without ar	y conditions, and					
rt II: (	(2) □ and/or	[II.2.8.3.2.	.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						
Pa	(2) □ and/or	[II.2.8.3.3.		the conditions referi Regulation (EU) 2020		pter 2, Sect	cion 1, point 6, of Annex V to		
	(2) □ and/or	[II.2.8.3.4.	•	the conditions referi Regulation (EU) 2020		pter 2, Sect	tion 1, point 7, of Annex V to		
	(2) □ and/or	[II.2.8.3.5.	•	the conditions referi Regulation (EU) 2020		pter 2, Sect	tion 1, point 8, of Annex V to		
	_			e 32(1), point (a), (b) down in Article 33 of		_	rated Regulation (EU) fulfilled.]]]]		
	II.3.		-	rledge and as declare bnormal mortalities	-		s come from establishments		
	II.4.	Arrangeme Regulation		-	nsignment in accor	dance with	n Article 4 of Delegated		
	II.5.		ea of anima	als, the period of val	-	_	. In the case of transport by extended by the duration of		
	(2)(6) □ [II.6.	establishme	ent approve		ations, none of the a	-	to the date of arrival to this the consignment has		
	(2) o either	[they come	from their	establishments of or	igin.]]				
(2) or [at least one of the animals of the consignment has undergone one assembly operation in an approved establishment.]]						bly operation in an			
	(2) or [at least one of the animals of the consignment has undergone two assembly operations in the approved establishments.]						bly operations in the		
	Animal welfa	ire attestatio	n						
			risions of Co				to be transported in urney due to start on		
	Notes								
	from the Euro Protocol on In	opean Union reland/North	and the Eu ern Ireland	ıropean Atomic Ene	rgy Community, and Annex 2 to that Pro	in particu otocol, refe	Britain and Northern Ireland lar Article 5(4) of the erences to the Union in this		
	provided for			e completed in accor to Commission Impl			ompletion of certificates 0/2235.		
	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.

Box "Place of destination": Indicate an establishment of the final destination of the consignment or an establishment approved for assembly operations in accordance with Articles 97 and 99 of Regulation

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	2024/1044 (2021/403) Model "OTHER-UNGULATES-INTRA-2
II. Health infor	rmation
I.12:	(EU) 2016/429.
Box reference I.17:	"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), based on which the animal health certificate for this consignment is issued in this establishment approved for assembly operations, may be indicated.
Box reference I.30:	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 reference I.30:	"Identification number": Indicate identification number of each animal.
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.
(5)	Insert the specific attestation(s) provided for in and required by the relevant legal act(s) adopted by the 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.
Stamp	

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