цО	KOPLAN	5111011			WODEL OV/CAP-INTRA-A			
	II. Health info	rmation						
	I, the unde	rsigned offi	cial veterinarian, hereby certify tha	at:				
	II.1.	The ovine/	caprine animals(1) of the consignm	ent described in Part I meet th	ne following requirements:			
		II.1.1.	They are identified as provided for in Article 45(2) or (4) or Article 46(1) of Commission Delegated Regulation (EU) 2019/2035.					
tion		II.1.2.	They, for at least the 30 day period if they are younger than 30 days o		consignment, or since birth,			
fica [.]			II.1.2.1. have been continuous	ly resident in the establishmer	nt of origin;			
Certi				act with kept ovine or caprine ovement restrictions for anima				
Part II: Certification				et or indirect contact with kept l country or territory during tl als.				
		II.1.3.	They have not shown clinical sign during the clinical examination w departure of the consignment, on	s or symptoms of diseases liste hich was carried out, within th datum van klinisch onderzoek invullen (insert date dd/n	ed for ovine/caprine animals ne 24 hour period prior to nm/yyyy).			
	II.2.	According requireme	to official information, the animals					
		II.2.1.	They do not come from establishin species or situated in a restricted a ovine/caprine animals.					
	(2)	either 🗹 [II.2.2.	They come from establishments fr B. suis without vaccination regard					
	(2)		either 🗹 [the establishments of or the status free from infection with ovine and caprine population;]					
	(2)		and/or □ [they have been subjected melitensis and B. suis with one of Commission Delegated Regulation sample taken during the 30 day per females taken at least 30 days after	the diagnostic methods provid (EU) 2020/688, carried out, wi eriod prior to departure, and i	led for in Part 1 of Annex I to th negative results, on a			
	(2)		and/or 🗆 [they are less than 6 mo	nths old;] N.V.T.				
	(2)		and/or 🗆 [they are castrated.] N.V.T)				
	(2)	or 0 [II.2.2.	They come from establishments fr B. suis with vaccination regarding Member State or zone thereof wit B. melitensis and B. suis regarding	; ovine and caprine animals ar hout the status free from infec	nd they are moved to a			
	(2)	either □ [II.2.3.	They are kept ovine animals and come from establishments in which infection with Mycobacterium tuberculosis complex (M. bovis, M. caprae and M. tuberculosis) has reported during the last 42 days prior to departure.] Anvinken indien de zending schapen bevat					
	(2)	and/or □ [II.2.3.	They are kept caprine animals and infection with Mycobacterium tub tuberculosis) has been carried out at least the 12 month period prior Regulation (EU) 2020/688.] Anivink	perculosis complex (M. bovis, M t on the caprine animals kept o	A. caprae and M. on the establishments during			
		II.2.4.	They come from establishments ir animals has not been reported du		-			
		II.2.5.	They come from establishments si establishments in which infection reported in kept animals of listed departure.	with epizootic haemorrhagic	disease virus has not been			
		II.2.6.	They come from establishments in the 15 day period prior to departu	-	nas not been reported during			

	II. Health infor	rmation						
		II.2.7.				which surra (Trypano eparture, and	osoma e	evansi) has not been reported
	(2)		either <mark>√</mark> [sı their depar		been repor	ted in the establishme	nts dur	ing the last 2 years prior to
ion	(2)							parture, following the last /ement restrictions until:
tificat				-	the infected and	l animals have been r	emoved	l from the establishments,
Part II: Certification				-	a test for su methods pr (EU) 2020/6	arra (Trypanosoma ev rovided for in part 3 of 88, carried out, with r aths after the infected hments.]	ansi) wi f Annex negative animals	nents have been subjected to ith one of the diagnostic I to Delegated Regulation results, on samples taken at s have been removed from
	(2)	□ [II.2.8.	They are ke	ept uncastra	ated male ov	ine animals, and Aan	vinken in bii mesto	dien zending rammen bevat. dieren!
			-				ididym	itis (Brucella ovis) has not
			_	carried out	t, with negat		le taker	oididymitis (Brucella ovis), I during the 30 day period I <mark>n te voldoen</mark>
	(2)	either □ [II.2.9. N.V.T.	(serotypes confirmed vaccinated 60 day peri	1-24), where during the l with a live od before tl	e no case of i last 24 mont vaccine agai he date of m	infection with blueton hs in the targeted anir nst infection with blu	gue vir nal pop etongue iremen	on with bluetongue virus us (serotypes 1-24) has been ulation and have not been e virus (serotypes 1-24) in the ts laid down in Article 020/688 are fulfilled.]
	(2)	and/or □ [II.2.9. N.V.T.	infection w	ith blueton	gue virus (se	erotypes 1-24) and the	require	adication programme for ements laid down in Article 020/688 are fulfilled, and
	(2)		either □ [II.2.9.1.	bluetongue	e virus (sero			y free from infection with h Article 40(3) of Commission
	(2)			either □ [II.2.9.1.1.	for at least	60 days prior to the da	ate of m	ovement]]
	(2)			and/or □ [II.2.9.1.2.	subjected to samples co animal into	o a serological test, wi llected at least 28 days	th nega follow zone se	ovement and have been tive results, carried out on ing the entry date of the asonally free from infection
	(2)			and/or □ [II.2.9.1.3.	subjected to collected at the Membe	o a PCR test, with nega	ntive res ng the en ally fre	ovement and have been sults, carried out on samples ntry date of the animal into e from infection with
	(2)		and/or □ [II.2.9.2.	place of de		d have been kept prot		uring transportation to the gainst attacks by vectors in a
	(2)			either □ [II.2.9.2.1.	for at least	60 days prior to the da	ate of m	ovement]]

	II. He	ealth information						
	(2)			and/or □ [II.2.9.2.2.	subjected to samples co		th negat s followi	
runcation	(2)				subjected to collected at	o a PCR test, with nega	ative res ng the da	ovement and have been ults, carried out on samples ate of the commencement of y vectors;]]]
Part II: Ceruncation	(2)	and/ [II.2.	9.3.	bluetongue State or zor	e virus whicl	n were reported durir vithin the immunity p	ig the pa	to 24 of infection with st 2 years in that Member aranteed in the
	(2)			either □ [II.2.9.3.1.	have been movement]	vaccinated more than]	60 days	before the date of
	(2)				PCR test, wi		l sample	l vaccine and subjected to a s collected at least 14 days specifications of the
	(2)	and/ [II.2.	9.4.	specific ant	tibodies agai		of infec	gical test able to detect tion with bluetongue virus e or zone and
	(2)			either □ [II.2.9.4.1.		ical test has been carr ore the date of mover		on samples collected at least
	(2)			and/or □ [II.2.9.4.2.	30 days bef subjected to	ore the date of the mo o a PCR test, with nega	ovement ative res	on samples collected at least and the animal has been ults, carried out on samples the date of movement;]]]
	(2)	[II.2.9. virus blue (c) o	s (serot tongue	ypes 1-24) r virus (sero	nor covered types 1-24) a	by the eradication pro	ogramm laid dow	n in Article 32(1)(a), (b) or
	(2)	II.2.9 en II.2.9.1 aanvinken indien dieren vanuit een tvbi worden gecertificeerd; bedrijf moet op de	9.1.	place of de		d have been kept prot		uring transportation to the ainst attacks by vectors in a
	(2)	"lijst tegen vector beschermo inrichtingen" staan (NVWA w	/ebsite).	either □ [II.2.9.1.1.	for at least	60 days prior to the d	ate of m	ovement]]
	(2)	1.1, 1.2 of 1.3 aan indien II.2.9 en II.2 aangevinkt, afhan wat van toepassing	vinken .9.1 is kelijk van	and/or □ [II.2.9.1.2.	subjected to samples col		th negat s followi	
	(2)			and/or □ [II.2.9.1.3.	subjected to collected at	o a PCR test, with nega	ative res ng the da	ovement and have been ults, carried out on samples ite of the commencement of y vectors;]]]
	(2)	and/ [II.2. II.2.9.2 Alleen mo bij aantonen immu middels serologie in combinatie met negatieve PCR te	9.2. gelijk unisatie	situated in establishme in Sections	a Member S ent, where s 1 and 2 of C	tate or in an area of a urveillance in compli	t least 1 ance wit Annex V	ure in an establishment 50 km radius centred on the h the requirements set out to Delegated Regulation d, and

II. Health information (2)the animals have been vaccinated against those serotypes from 1 to either 🗆 [II.2.9.2.1. 24 of infection with bluetongue virus which were reported during the past 2 years in an area of at least 150 km radius centred on the place where the animals were kept and are within the immunity N.V.T. period guaranteed in the specifications of the vaccine and (2) have been vaccinated more than 60 days before the date either 🗆 Part II: Certification [II.2.9.2.1. of movement]]] N.V.T. 1 (2) and/or 🗆 have been vaccinated with an inactivated vaccine and [II.2.9.2.1. subjected to a PCR test, with negative results on samples N.V.T. 2. collected at least 14 days after the onset of the immunity set in the specifications of the vaccine;]]]] (2) and/or □ the animals have been immunised against those serotypes from 1 to II.2.9.2.2 aanvinken [II.2.9.2.2. 24 of infection with bluetongue virus which were reported during als dmv serologie is aangetoond the past 2 years in an area of at least 150 km radius centred on the dat de dieren antilichamen hebber tegen BT-3 in combinatie met place where the animals were kept, and een negatieve PCR test (2) either 🗆 the animals have been subjected with positive results to a [II.2.9.2.2. N.V.T. 60 days before the date of movement]]] 1. (2) and/or □ [II.2.9.2.2. II 2 9 2 2 2 aanvinken als dmv serologie is aangetoond dat 2 de dieren antilichamen hebben tegen BT-3 in combinatie met een negatieve PCR test movement:1111 (2) and/or □ They do not fulfil the requirements laid down in points 1 to 3 of Section 1 of Chapter 2 of [II.2.9. Part II of Annex V to Delegated Regulation (EU) 2020/689 and the competent authority of the 2.9 en II.2.9.1 aanvinken als (12.5 cf 11.2.5 cf administration in the semming vrij is van BT én vermeld staat in de EU lijst met derogaties (2) én als aan de in de derogatie genoemde eisen wordt voldaan Zie BT-07 voor de link naar de derogaties. zone thereof either \Box [II.2.9.1. and (2)either 🗆 [II.2.9.1.1. Delegated Regulation, and N.V.T. (2) II.2.9.1.2 aanvinken als and/or □ lidstaat van bestemming vrij is van BT en er middels derogatie [II.2.9.1.2. Delegated Regulation, and wordt gecertificeerd. (2) and/or □ [II.2.9.1.3. Delegated Regulation, and N.V.T. and/or \Box (2)[II.2.9.1.4. Delegated Regulation, and N.V.T. BT dient het VVM behandeld te worden. Delegated Regulation are fulfilled.]]] (2)and/or □ [II.2.9.2. (serotypes 1-24) and the Member State of destination has informed the Alleen aanvinken indien dieren

serological test carried out on samples collected at least

the animals have been subjected with positive results to a serological test carried out on samples collected at least 30 days before the date of the movement and to a PCR test, with negative results, carried out on samples collected not earlier than 14 days before the date of

Member State of origin authorised movement of those animals to another Member State or 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)(a), (b) and (c) of Delegated Regulation (EU) 2020/689,

point 5 of Section 1 of Chapter 2 of Part II of Annex V to that

point 6 of Section 1 of Chapter 2 of Part II of Annex V to that

point 7 of Section 1 of Chapter 2 of Part II of Annex V to that

point 8 of Section 1 of Chapter 2 of Part II of Annex V to that

Bij vervoer DOOR een lidstaat the requirements laid down in Article 32(1)(a), (b) or (c) van of met uitr.progr. voor or Article 32(2) of Delegated Regulation (EU) 2020/688 and the requirements laid down in Article 33 of that

with an approved eradication program for infection with bluetongue virus Commission and the other Member States that such movement is authorised naar deel van Spanie gaan met uitroeiingsprogramma, en als aan de eisen zoals vermeld in de under the conditions referred to in Article 43(2)(a), (b) and (c) of Delegated anse derogatie is voldaan. Regulation (EU) 2020/689, and

(2)

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	II. Hea	alth information				
	(2)	Indien gebruik gemaakt wordt va de Spaanse derogatie en dieren gaan naar gebied met uitroeiingsprogramma, dient II.2.9.2.2 aangevinkt	and/or □ [II.2.9.2.2.	-	ection 1 of Chapter 2 of Part . Regulation, and	II of Annex V to that
	(2)	II.2.9.2.2 aangevinkt te worden.	and/or □ [II.2.9.2.3.	point 7 of S	ection 1 of Chapter 2 of Part . Regulation, and <mark>N.V.T.</mark>	II of Annex V to that
ion	(2)		and/or □ [II.2.9.2.4.	-	ection 1 of Chapter 2 of Part Regulation, and <mark>N.V.T.</mark>	II of Annex V to that
ertificat		Bij vervoer DO vrij van of met BT dient het V behandeld te v	uitr.progr. voor /M	32(2) of Del	ments laid down in Article 3 egated Regulation (EU) 2020/ n Article 33 of that Delegated	688 and the requirements
Part II: Certification	(2)	Alleen aanvinken indien land van bestemming beperkingsgebied is en dat land onder derogatie voorwaarden dieren accepteert.	by the erac 24) and the	dication prog e Member St	tion with bluetongue virus (s gramme for infection with bl ate of destination has inform hat such movement is author	uetongue virus (serotypes 1- ed the Commission and the
	(2)		either □ [II.2.9.3.1.	without an	y conditions, and N.V.T.	
	(2)		and/or □ [II.2.9.3.2.	2 of Part II	of Annex V to Delegated Regi	ooint 5 of Section 1 of Chapter Ilation (EU) 2020/689, and N.V T.
	(2)	Voor derogaties waarbij bescherming en PCR test gevraagd wordt, dient II.2.9.3. aangevinkt te worden.		of Part II of	conditions referred to in poin Annex V to Delegated Regula	ation (EU) 2020/689, and
	(2)	Voor derogaties van België er	and/or □ [II.2.9.3.4.	of Part II of		ation (EU) 2020/689, and N.V.T
	(2)	deelstaten van Duitsland (waa alleen BTV3 heerst), dient II.2.9.3.5 aangevinkt te worde	^{1.} [II.2.9.3.5.	of Part II of	onditions referred to in poin Annex V to Delegated Regula	ation (EU) 2020/689, and
		Bij vervoer DO vrij van of met BT dient het V behandeld te v	uitr.progr. voor √M	32(2) of Del	ments laid down in Article 3 egated Regulation (EU) 2020/ n Article 33 of that Delegated	688 and the requirements
	(2)	[II.2.10. of Section Parliamer Member S	A of Chapter at and of the tate listed in	r A of Annex Council as h point 3.2. of	mber State or zone of a Mem VIII to Regulation (EC) No 99 aving a negligible risk status that Section as having an ap anninken indien dieren gaan na	9/2001 of the European for classical scrapie or for a proved national scrapie
	(2)	either o [u in point 2.	come from a 3. of Section	holding situ	ated in a Member State or zo r A of Annex VIII to Regulation	
	(2)	accordano 999/2001 a	e with point and listed as	1.2. of Sections such by the	cognised as having a negligib on A of Chapter A of Annex V competent authority of the M vinken indien primair bedrijf op lijst ieren gaan naar Aut, Cze, Fin, Sw	III to Regulation (EC) No
	(2)	and/or □ Chapter B species ar	[come from a of Annex VI d are of the	a holding no I to Regulatio ARR/ARR pri	t subject to the measures laid on (EC) No 999/2001 and the a on protein genotype, or the a	down in points 3 and 4 of animals are of the ovine
	(2)				ned for an approved body, in ve 92/65/EEC.]	nstitute or centre as defined
	(2)				et out in point 4.1.(d) of Section. .]] <mark>Alléén van toepassing bij zeldz</mark>	on A of Chapter A of Annex ame rassen. Zie verder K-LV-SGIU- <mark>01</mark>
	(2)	[II.2.10. State othe Regulation than those	 or The animals are for breeding and are intended for a Member State or zone of a Member [II.2.10. State other than those listed in point 2.3. of Section A of Chapter A of Annex VIII to Regulation (EC) No 999/2001 as having a negligible risk status for classical scrapie or other than those listed in point 3.2. of that Section as having an approved national scrapie control 			
		programm	re, unu <mark>Adily</mark>		n naar andere lidstaat gaan dan Au	, 525, F III, 6W6, BIIK OF 6VII

either \circ [come from a holding situated in a Member State or zone of a Member State listed in point 2.3. of Section A of Chapter A of Annex VIII to Regulation (EC) No 999/2001 as having a negligible risk status for classical scrapie.] and/or \Box [come from a holding recognised as having a negligible risk of classical scrapie in accordance with point 1.2. of Section A of Chapter A of Annex VIII to Regulation (EC) No 999/2001 and listed as such by the competent authority of the Member State in accordance with point 1.1. of that Section.] Any intercontraction of the high primair bedrijf op lijst verwaarloosbaar risico scrapie staat en dieren gaan naar andere lidstaat dan Aut, Cze, Fin, Swe, Dnk of Svn and/or \Box [come from a holding recognised as having a controlled risk of classical scrapie in accordance with point 1.3. of Section A of Chapter A of Annex VIII to Regulation (EC) No 999/2001 and listed as such by the competent authority of the Member State in accordance with point 1.1. of that Section.] Anvinken indien primair bedrijf op lijst gecontroleerd risico scrapie staat en dieren gaan naar andere lidstaat dan Aut, Cze, Fin, Swe, Dnk of Svn and/or \Box [come from a holding not subject to the measures laid down in points 3 and 4 of Chapter B of Annex VII to Regulation (EC) No 999/2001 and the animals are of the ovine species and are of the ARR/ARR prion protein genotype, or the animals are of the caprine species and carry at least one of the K222, D146 or S146 alleles.] and/or [] [come from and are destined for an approved body, institute or centre as defined in Article 2(1)(c) of Directive 92/65/EEC.1 or \circ [comply with the conditions set out in point 4.1.(d) of Section A of Chapter A of Annex VIII to Regulation (EC) No 999/2001.]] Alléén van toepassing bij zeldzame rassen. Zie verder K-LV-SGIU-01 The animals are not for breeding and are intended for a Member State or zone of a Member or o [II.2.10. State other than those listed in point 2.3. of Section A of Chapter A of Annex VIII to Regulation (EC) No 999/2001 as having a negligible risk status for classical scrapie or other than those listed in point 3.2. of that Section as having an approved national scrapie control Aanvinken indien het om mestdieren gaat én lidstaat van bestemming anders is dan Aut, Cze, Fin, Swe, Dnk of Svn programme.] To the best of my knowledge and as declared by the operator, the animals come from establishments where there were no abnormal mortalities with an undetermined cause. (2) [II.4. According to official information and as declared by the operator, they are semen donor animals, and II.4.1. they come from a semen collection centre and will be transported directly to another semen collection centre in accordance with Article 19 of Commission Delegated Regulation (EU) 2020/686; and either \circ they were continuously present since the date of their admission at the semen collection [II.4.2. centre and were subjected, with negative results, to all compulsory routine tests referred to in point 2 of Chapter I of Part 3 of Annex II to Delegated Regulation (EU) 2020/686 in the period of the preceding 12 months prior to date of that movement; and] they were subjected, with negative results, to all tests referred to in point 1(c) and (d) of or o [II.4.2. Chapter I of Part 3 of Annex II to Delegated Regulation (EU) 2020/686, required before admission to a semen collection centre carried out during the period immediately preceding quarantine and during the guarantine period; and] II.4.3. the prior consent of the centre veterinarian of the semen collection centre of destination has been obtained by the operator; and the means of transport used have been cleansed and disinfected before use.] II.4.4. Arrangements are made to transport the consignment in accordance with Article 4 of Delegated Regulation (EU) 2020/688. This certificate is valid for 10 days from the date of issuing. In the case of transport by waterway/sea of animals, the period of validity of the certificate may be extended by the duration of the journey by waterway/sea. (2)(3) Since leaving their establishments of origin and before arriving to this establishment approved for assembly operations, none of the animals of the consignment has undergone more than two assembly vierkantje aanvinken ndien gecertificeer@perations, and ordt vanaf verzam either \circ [they come from their establishments of origin.]] or \circ [at least one of the animals of the consignment has undergone one assembly operation

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II.3.

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II.5.

II.6.

[II.7.

(2)

(2)

Part II: Certification

II. Health information

(2)	(2) or ○ [at least one of the animals of the consignment has undergone two assembly operations on approved establishments.]] Dit bolletje aanvinken indien minstens 1 vd dieren afkomstig is van een ander Nederlands verzameter									
Anim	Animal welfare attestation									
At the time of inspection, the animals covered by this 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) (4)(5).										
Notes	s:									
In acc from	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 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									
	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	I:									
Box refere I.11:	ence	"Place of dispatch": Indicate an establishment of the origin of the animals in the consignment or an e 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.								
Box refere I.12:	ence	"Place of destination": Indicate an establishmen establishment approved for assembly operations (EU) 2016/429.								
Box refere I.17:	ence	"Accompanying documents": In case the animals are dispatched from an establishment approved for e 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.								
	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.									
Box refere I.30:	ence	"Identification number": Indicate identification codes of the animals in the consignment identified in accordance with Article 45(2) or (4) or Article 46(1) of Delegated Regulation (EU) 2019/2035.								
Part I	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.								
(4) This statement does not exempt transporters from their obligation in accordance with Union rule force in particular regarding the fitness to be transported.										
(5)	ring Offi	To be completed in case of consignment grouped located in the Member State of transit. icer/Official veterinarian	l in an establishment appro	ved for assembly operations						
-	-		Dualification and title							
Date o	of signa p	-	ignature							