_								
	I.1. Consignor				I.2. IMSOC reference		I.2.a. Local refere	ence
	Name						I.3. Central Comp	netent Authority
	Address		****					
	Country		ISO Co	de			I.4. Local Compe	tent Authority
nt	I.5. Consignee				I.6. Operator conducting	g assembly op	perations indepen	dently of an
ot consignment	Name				establishment			
3	Address				Name			
ĭ	Country		ISO Co	de	Address Approval			
51					Number			
ı					Country		ISO Code	
	I.7. Country of orig	σin		ISO Code	I.9. Country of destinati	on		ISO Code
	in country of one	5***			list country of destinati	011		
Į.	I.8. Region of origi	in		Code	I.10. Region of destinati	on		Code
Part I: Description								
al E	I.11. Place of dispa	atch			I.12. Place of destination	ı		
::	Name				Name			
11	Address Approval				Address			
ية الت	Number				Approval Number			
	Country		ISO Co	de	Country		ISO Code	
	I.13. Place of loadi	ing			I.14. Date and time of de	nartura		
	Name	ing			1.14. Date and time of ut	eparture		
	Address							
	Approval							
	Number		ISO Co	do				
	Country		150 00	ue				
	I.15. Means of Tra	nsport	1		I.16. Transporter			
	Mode	International transport	Identificati	on	Name			
		document			Address			
					Approval Number			
					Country		ISO Code	
					I.17. Accompanying doc	uments		
		1			Document Type Accompanying docume	n+		
					reference	:111		
					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							
	Germinal product	s 🗆						
	I.21. For transit th	rough a third coun	try					
	Third country				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	
	I.24. Estimated jou	ırney time		ı	I.25. Journey Log	T		
	I.26. Total number			I.27. Total quantity		I.28. Total g	ross weight	
- 1	I.30. Description o	_						
	1. 05 PRODUCTS O	OF ANIMAL ORIGIN	, NOT ELSE	WHERE SPECIFIED OR II	ICLUDED			
	0511 Animal pro 051199 Other		ere specified	or included; dead anima	als of Chapter 1 or 3, unfi	t for human o	consumption	
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	NION					IK
- 1	#1. Commodity	Identification Number	Quantity	Nature of commodity	Identification Mark	
	Species	Package count	Date of collection / production	Plant / Establishment / Centre	Туре	
5						
Tart I Took I bridge or county						
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	II. Health info	rmation						
	I, the unde	rsigned offi	cial veterina	arian, hereb	y certify tha	t:		
		_			-	animals describe	d in Part I:	
uc			II.1.1.	has been c	ollected, pro	cessed and store	d, and dispat	ched from the semen register by the competent
Part II: Certification			II.1.2.	collection operationa	centre which al procedure	complies with r	equirements quipment set	ched from the semen as regards responsibilities, out in Part 1 of Annex I to
Part II:			(1) either	○ [II.1.3.	to moveme established diseases su or those res collected be	nt restrictions af for reasons of li bject to emergen strictions do not efore the restrict	fecting ovine sted diseases cy measures apply to this ions were est	centre or a zone not subject and caprine animals and relevant for those species or relevant for those species, semen because it was ablished, and has not been ealth status for an adequate
<u> </u>			(1) or	○ [II.1.3.	movement and establi	restrictions affe	cting ovine ar (3), bu	centre or a zone subject to nd caprine animals species t derogations from d, and:
			(1)	☐ [it com	plies with the	e requirements s	et out in	(4);]]]
			(1)	☐ [and in	particular, i	t is	(5).]]]	
	(1)	[II.1.			ne] (1) 🏻 [c and dispatch		als described	l in Part I has been collected,
			(1) either	○ [II.1.1.	affecting or listed disea emergency	vine and caprine ses relevant for t measures releva	animals and hose species ant for those	o movement restrictions established for reasons of or diseases subject to species, and has not been in the status for an adequate
			(1) or	○ [II.1.1.		rine and caprine (3), but derog	animals spec	ovement restrictions cies and established for novement restrictions have
			(1)	☐ [it com	plies with the	e requirements s	et out in	(4);]]
			(1)	☐ [and in	particular, i	t is	(5).]]	
			II.1.2.			re the donor anii n (EU) 2020/686, a		as referred to in Article 13
				II.1.2.1.	_	_		of the competent authority cept the consignment;
				II.1.2.2.		nimals have bee date of semen c	-	xamined by a veterinarian
				II.1.2.3.	least the in	_	ded for in Art	shment which include at icle 8(1), point (a), of
				(1) ○ either	[II.1.2.4.	described in Par have been kept holdings recogn risk of classical point 1, of Anne	rt I was collect continuously ised as havin scrapie accor x VIII to Regu	☐ [caprine] (1) animals ted from animals which since birth on a holding or g a negligible or controlled ding to Chapter A, Section A, alation (EC) No 999/2001 of d of the Council, except

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	II. Health in	formation					
						during the period when they collection centre that compli the conditions set out in Chap 1.3(c)(iv), of Annex VIII to Re	ed during that period with pter A, Section A, point
Part II: Certification				(1) ○ or	[II.1.2.4.	the semen of \square [ovine] (1) described in Part I was collected have been kept continuously the collection on a holding of complied for the last three years with the requirements laid depoints 1.3(a) to (f), of Annex 1999/2001, except during the part as semen collection centred at a semen collection centred period with the conditions see A, point 1.3(c)(iv), of Annex 1999/2001.]	cted from animals which for the last 3 years before holdings which has/have ears before the collection own in Chapter A, Section A, WIII to Regulation (EC) No period when they were kept that complied during that et out in Chapter A, Section
	_			(1) ∘ or	[II.1.2.4.	the semen of \square [ovine] (1) described in Part I was collected have been kept continuously State or zone of a Member St Section A, point 2.3, of Annex 999/2001 as having a negligible scrapie;]	cted from animals which since birth in a Member ate listed in Chapter A, x VIII to Regulation (EC) No
				(1) ○ or	[II.1.2.4.	the semen of ovine animals of collected from ovine animals protein genotype;]	
	II.2.	The seme		in Part I is i	ntended for	artificial reproduction and wa	s obtained from donor
			II.2.1.			emained since birth in the Uni vith the requirements for entr	
			II.2.2.	II.2.6, fron establishn	n establishm nents under	of commencement of the quar nents in a Member State or zon official control by the compete r a zone thereof:	e thereof, or from
				II.2.2.1.	reported v least 30 da	an area where foot -and -mou vithin a 10-km radius centred o lys and in which foot and mou luring at least 3 months, and:	on the establishment for at
		(1)	o either	[the donor	r animals we	ere not vaccinated against foot	and mouth disease;]
				(1)	or	[the donor animals were vac mouth disease during 12 mor collection of the semen but n immediately prior to the date and 5 % (with a minimum of quantity of semen taken fron time is submitted to a virus is mouth disease with negative	nths prior to the date of not during the last 30 days e of collection of the semen, five straws) of each in the donor animals at any solation test for foot and
				II.2.2.2.	and the do	infection with Brucella abortu onor animals have never been nent of a lower health status;	
				(1) (6)	□ [II.2.2.3.	in which infection with Myco complex (M. bovis, M. caprae not been reported during the	e and M. tuberculosis) has
				(1) (7)	□ [II.2.2.3.	in which surveillance for infetuberculosis complex (M. box	-

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UI	NION				3024/1044 (2021/403) MOD	EL OV/CAP-SEM-A-INTRA
	II. Health information					
Part II: Certification					tuberculosis) has been carried animals kept in the establish months, as referred to in Art Delegated Regulation (EU) 20 this period, infection with M complex (M. bovis, M. capraed been reported in caprine ani establishment, measures were Part 1, point 3, of Annex II to	ments during at least 12 icle 15(3) of Commission 020/688, and in case, during ycobacterium tuberculosis and M. tuberculosis) has imals kept in the re taken in accordance with
II: Cer			II.2.2.4.	in which su the last 30 c	ırra (Trypanosoma evansi) ha days, and:	ns not been reported during
Part		(1)	o either	[surra has i years;]	not been reported in the estab	olishments during the last 2
		(1)	or	years and f establishme the date on establishme have been s methods pr (EU) 2020/6 least 6 mon	been reported in the establish following the date of the last of ents have remained under mowhich the infected animals hents, and the remaining anim subjected to a test for surra we rovided for in Part 3 of Annex (88, carried out, with negative of this following the date on white removed from the establishm	outbreak, the affected ovement restrictions until have been removed from the hals in the establishments with one of the diagnostic of I to Delegated Regulation e results, on samples taken at lich the infected animals
			(1) (6)		in which ovine epididymitis reported during the last 12 m	
			(1) (11)	□ [II.2.2.6.	where, during the last 60 day quarantine accommodation of they have been subjected, wi serological test for ovine epid any other test for ovine epid equivalent documented sens required in accordance with of Annex II to Delegated Region	referred to in point II.2.6, ith negative results, to a didymitis (Brucella ovis), or idymitis (Brucella ovis) of an itivity and specificity, as Part 3, Chapter I, point 1(b),
		II.2.3.		ir admission	s or clinical signs of transmis to a semen collection centre a	
		II.2.4.		-	fied as provided for in Article ed Regulation (EU) 2019/2035	
		II.2.5.		t 30 days priction period:	or to the date of first collection	n of the semen and during
			II.2.5.1.	movement mouth dise Valley fever sheep pox a	n establishments situated in a restrictions established due to ase, infection with rinderpest r virus, infection with peste d and goat pox or contagious ca ging disease relevant for ovin	o the occurrence of foot and t virus, infection with Rift es petits ruminants virus, prine pleuropneumonia, or
			II.2.5.2.	abortus, B. tuberculosi rabies, anth epizootic ha virus (serot caprine ani	n a single establishment when melitensis and B. suis, infection is complex (M. bovis, M. capra nrax, surra (Trypanosoma eva aemorrhagic disease virus, in trypes 1-24) and, in case of oving imals which are kept together is (Brucella ovis) have not bee	on with Mycobacterium he and M. tuberculosis), hensi), infection with fection with bluetongue he animals and those with ovine animals, ovine
			II.2.5.3.	were not in	contact with animals from e	stablishments situated in a
	<u> </u>					

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	NION				2024/1044 (2021/403) MODEL OV/CAP-SEM-A-INTRA
	II. Health information				
				diseases re	ct to movement restrictions due to the occurrence of eferred to in point II.2.5.1 or from establishments which do ne conditions referred to in point II.2.5.2;
			II.2.5.4.	were not u	sed for natural breeding;
Part II: Certification		II.2.6.	accommod health stati	ation, wher us were pre	o a quarantine for at least 28 days in quarantine e only other cloven-hoofed animals with at least the same sent, which on the day of admission of the donor animals a centre complied with the following conditions:
t II: Cer			II.2.6.1.	established	ated in a zone not subject to movement restrictions due to diseases referred to in point II.2.5.1 or it was ogation as referred to in point II.1.1, if applicable;
Par			II.2.6.2.	none of the	e diseases referred to in point II.2.5.2. has been reported 30 days;
			II.2.6.3.	reported w	ated in an area where foot and mouth disease has not been within a 10-km radius centred on the quarantine lation for at least 30 days;
			II.2.6.4.	least 3 mor	no outbreak of foot and mouth disease reported during at nths preceding the date of admission of the donor animals men collection centre;
		II.2.7.	were kept i	in the semer	n collection centre:
			II.2.7.1.	established	situated in a zone not subject to movement restrictions due to diseases referred to in point II.2.5.1 or it was ogation as referred to in point II.1.1, if applicable;
			II.2.7.2.		e of the diseases referred to in point II.2.5.2 has been or at least 30 days prior to the date of collection of the d:
			(1) (6) either	o [at least 3	30 days following the date of the collection;]
			(1) (7) or		e date of departure of the consignment of semen to ember State;]
			II.2.7.3.	reported w	an area where foot and mouth disease has not been within a 10-km radius centred on the semen collection at least 30 days; and:
			(1)(6) either		m foot and mouth disease for at least 3 months prior to the lection of the semen and 30 days from the date of
			(1)(7) or	date of coll consignme animals ha continuous	m foot and mouth disease for at least 3 months prior to the lection of the semen and until the date of dispatch of the nt of semen to another Member State and the donor are been kept at that semen collection centre for a seperiod of at least 30 days immediately prior to the date of of the semen;]
			th at least or types 1-24):	ne of the foll	lowing conditions as regards infection with bluetongue
		(1)	o either	[II.2.8.1.	they have been kept for at least 60 days prior to and during collection of the semen in a Member State or zone thereof free from infection with bluetongue virus (serotypes 1-24) where no case of infection with bluetongue virus (serotypes 1-24) in the targeted animal population has been confirmed during the last 24 months;]
		(1)	o or	[II.2.8.2.	they have been kept in a seasonally disease-free zone, during the seasonally disease-free period, for at least 60

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	II. Health information						
					Member S eradicatio	tate or zone there	lection of the semen, in a of with an approved inst infection with 1-24);]
n H		(1)	or	[II.2.8.3.	during the days prior Member S authority semen has competent to the cond	to and during coltate or zone there of the place of origonatined the price authority of the Iditions for establish	ssonally disease-free zone, se-free period, for at least 60 lection of the semen, in a of where the competent gin of the consignment of or written consent of the Member State of destination shment of that seasonally ept the consignment of
		(1)	□ and/or	[II.2.8.4.			ctor protected establishment and during collection of the
		(1)	□ and/or	[II.2.8.5.	antibodies negative r	to the bluetongue	a serological test to detect e virus serotypes 1-24, with and 60 days from the date of c;]
		(1)	□ and/or	[II.2.8.6.	for bluetor results, on final collections semen at i	ngue virus (seroty blood samples ta ction of the semen ntervals of at leas solation test, or of	an agent identification test pes 1-24), with negative ken at commencement and and during collection of the t every 7 days, in the case of at least every 28 days, in the
	II.2.9.		vith at least o hagic disease		_	litions as regards	infection with epizootic
			(1)	o either	[II.2.9.1.	to and during co Member State on has not been rep	kept for at least 60 days prior llection of the semen in a zone thereof where EHDV ported within a radius of 150 ishment for at least 2 years;]
			(1)	o or	[II.2.9.2.	free zone, during	kept in a seasonally diseaseg the seasonally disease-free list 60 days prior to and n of the semen;]
			(1)	□ and/or	[II.2.9.3.	establishment fo	kept in a vector protected or at least 60 days prior to ction of the semen;]
			(1) or	o [II.2.9.4.	official fin	dings the followin and have been	er State in which according to ng serotypes of EHDV exist: n subjected with negative
						each case to the fo laboratory:	llowing tests carried out in
				(1)	o either	antik resu thro and the c	rological test to detect codies to EHDV, with negative lts, at least every 60 days ughout the collection period between 28 and 60 days from late of the final collection of emen;]]
				(1) 🗆	[II.2.9.4.2.	an agent identifi	cation test for EHDV, with

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	II. Health information					
on				and/or	the commenceme semen and durin at intervals of at	on blood samples taken at ent and final collection of the g the collection of the semen least every 7 days, in the ation test, or of at least every se of PCR.]
Part II: Certification		(1) (8)	□ [II.2.10.	taken with quarantine in accordan	subjected to the following test in 30 days prior to the date of referred to in point II.2.6, wince with Part 3, Chapter I, poince with Part 3, Chapter I, poince with CEU) 2020/686:	commencement of the th negative results, required
Part			II.2.10.1.	serological	n with Brucella abortus, B. m test referred to in Part 1, poir (EU) 2020/688;	
			(1) (11)	□ [II.2.10.2.	for ovine epididymitis (Bruce any other test with an equiva and specificity;]	_
	II.2.11.	after the d results, red	ate of comm	encement o cordance wi	ng tests, carried out on sample f the quarantine referred to in th Part 3, Chapter I, point 1(d)	n point II.2.6, with negative
			II.2.11.1.	serological	n with Brucella abortus, B. m test referred to in Part 1, poir (EU) 2020/688;	
			(1) (11)	□ [II.2.11.2.	for ovine epididymitis (Bruce any other test with an equiva and specificity;]	
		II.2.12.	following o	compulsory	semen collection centre, at le routine tests, required in acco o Delegated Regulation (EU) 2	ordance with Part 3, Chapter
			II.2.12.1.	serological	n with Brucella abortus, B. m test referred to in Part 1, poir (EU) 2020/688;	
			(1) (11)	□ [II.2.12.2.	for ovine epididymitis (Bruce any other test with an equiva and specificity.]	ella ovis), a serological test or alent documented sensitivity
			(1) (12)	□ [II.2.13.	have been subjected to the following blood samples taken within 3 the semen, with negative res	30 days prior to collection of
			II.2.13.1.	serological	n with Brucella abortus, B. m test referred to in Part 1, poir (EU) 2020/688;	
			(1) (11)	□ [II.2.13.2.	for ovine epididymitis (Bruce any other test with an equive and specificity;]]	ella ovis), a serological test or alent documented sensitivity
	II.3.	The semen	described i	n Part I		
		(1) (8)	□ [II.3.1.	health requ	ollected, processed and stored airements set out in Part 1, po Regulation (EU) 2020/686;]	
		II.3.2.	accordance	e with requi	other packages on which the n rements provided for in Artic mark is indicated in box I.30;	le 10 of Delegated Regulation
		II.3.3.	is transpor	ted in a con	tainer which:	

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	II. Health info	rmation							
				II.3.3.1.	semen col veterinari	l and numbered particular to the lection centre und an, or by an official indicated in box	er responsil al veterinari	oility of the centi	re
ion				II.3.3.2.		leaned and either container;	disinfected	or sterilised bef	ore use, or is
ificat				(1) (9)	□ [II.3.3.3.	has been filled ir been previously			ich has not
Part II: Certification		(1) (13)	□ [II.4.	Where an in Part I:	antibiotic o	r mixture of antibi	otics was ac	lded to the seme	n described
Part I			II.4.1.			ic or mixture of an is contained in the			the semen (14).
	II.4.2.	was kept	at a tempera	ature of at le	east 5°C for r	ic(s), and before a ot less than 45 min cidal activity.]			
	Notes:								
			_			he United Kingdo			
	Protocol or	n Ireland/N	orthern Irel	and in conj	unction with	gy Community, an Annex 2 to that P espect of Northeri	rotocol, refe		
				_		dance with the no ementing Regulati		-	rtificates
	Part I:								
	Box reference I.11:	collection 2020/686,	centre or, ii	n case of an registration	establishme	oval number and t nt referred to in A d address of the es	rticle 13 of	Delegated Regula	ation (EU)
	Box reference I.12:					nd unique registra ent of semen.	tion or appr	oval number of	the
	Box reference I.19:	Seal num	ber shall be	indicated.					
	Box reference I.26:	Total nun	nber of pack	ages shall c	orrespond to	the number of co	ntainers.		
	Box reference I.30:	"Species":	: Select amoi	ngst "Ovis a	ries" or "Cap	ra hircus" as appr	opriate.		
	"Type": Inc	dicate seme	en.						
	"Identifica	tion numb	er": Indicate	the identifi	ication numl	oer of each donor a	animal.		
			ation mark" ent is placed		ie mark on t	ne straw or other p	oackages wh	ere the semen o	of the
		"Date of c	collection/pro	oduction": I	ndicate the o	late on which the	semen of the	e consignment w	vas collected.
		of the sen Delegated	nen collectio	n centre or, (EU) 2020/6	, in the case 86, the uniq	ablishment/centre of an establishmer ue registration nur	it as referre	d to in Article 13	3 of
		"Quantity	": Indicate t	he number	of straws or	other packages wi	th the same	mark.	
	"Test": Ind	icate for B	ΓV-test: poin	t II.2.8.5 and	d/or point II.	2.8.6, and/or for EI	HD-test: poir	nt II.2.9.4.1 and/o	or point

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2024/1044 (2021/403) MODEL OV/CAP-SEM-A-INTRA

Health information
Part II: (1) Delete if not applicable. (2) Only semen collection centres approved by the competent authority and included in the register referred to in Article 101(1), point (b), of Regulation (EU) 2016/429 of the European Parliament and of the Council and Article 7 of Delegated Regulation (EU) 2020/686. (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 159(2), points (a), (b) and (c), of Regulation (EU) 2016/429. (6) Applicable for caprine animals. (7) Applicable for caprine animals. (8) Applicable for semen collected at a semen collection centre. (9) Applicable for frozen semen. (10) Applicable for ovine animals and for those caprine animals which are kept together with ovine animals and for those caprine animals which are kept together with ovine animals (12) Applicable for semen collected at an establishment where the donor animals are kept as referred to in Article 13 of Delegated Regulation (EU) 2020/686. (13) Mandatory attestation in case antibiotic(s) was/were added. (14) Insert the name(s) of the antibiotic(s) added and its (their) concentration or the commercial name of the semen diluent containing antibiotic(s). Certifying Officer/Official veterinarian Name (in capital letters) Oqualification and title Signature
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