INTRA

	I.1. Consignor				I.2. IMSOC reference		I.2.a. Local refere	ence
	Name Address						I.3. Central Comp	etent Authority
	Country		ISO Code				I.4. Local Compe	tent Authority
ent	I.5. Consignee				I.6. Operator conducting establishment	g assembly o _l	perations indepen	dently of an
u	Name Address				Name			
\mathbf{sig}	Country		ISO Code		Address			
ons					Approval Number			
$^{ m c}$					Country		ISO Code	
Part I: Description of consignment	I.7. Country of ori	gin	ISO Code	!	I.9. Country of destination	on		ISO Code
crip	I.8. Region of original	in	Code		I.10. Region of destination	on		Code
)es	I.11. Place of dispa	atch			I.12. Place of destination	າ		
I: I	Name				Name			
ırt	Address Approval				Annous			
Ρĉ	Number				Approval Number			
	Country		ISO Code		Country		ISO Code	
	I.13. Place of loadi	ing			I.14. Date and time of de	eparture		
	Name							
_	Approval							
	Approval Number							
	Country		ISO Code					
	I.15. Means of Tra		- 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		I.16. Transporter			
	Mode	International transport	Identification		Name			
		document			Address Approval			
					Number		100 0 1	
					Country		ISO Code	
					I.17. Accompanying doc	uments		
					Document Type Accompanying docume	nt		
					reference	:111		
					Date of Issue			
					Country Place of issue			
	I.18. Transport co	nditions						
	Chilled		Ambient 🗆			Frozen 🗆		
	I.19. Container No	/ Seal No						
	I.20. Certified as							
	Germinal product	s L						
	I.21. For transit th	rough a third coun	try					
	Third country				ISO Code			
	Exit point Entry point				BCP code BCP code			
		rough Member Sta	te(s)		I.23. For export			
	Member State	5	ISO Code		Third country		ISO Code	
	I.24. Estimated jou	ırney time			Exit point I.25. Journey Log		BCP code	
	I.26. Total number		I.27. Total quan	ntity		I.28. Total g	ross weight	
	I.30. Description o		1			1	- 0 ,	
	_	_	, NOT ELSEWHERE SPECIFIE	D OR INC	CLUDED			
		oducts not elsewhe	re specified or included; dea			t for human (consumption	
	05119985 (Other						

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_	ION	1	T	T		INIK
	#1. Commodity	Identification Number	Quantity	Nature of commodity	Identification Mark	
	Species	Package count	Date of collection / production	Plant / Establishment / Centre	Туре	
5						
t at a transfer of country						
b						
1						
I						
I						
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	II. Health info	rmation				
	I the unde	reigned offi	cial veterinari	ian, hereby certify tha	t ·	
		_			os] (1) of porcine animals des	scribed in Part I have been
	(1) = [11.1.				d by the embryo collection tea	
		II.1.1	is approved a	and kept in a register l	by the competent authority;	
Part II: Certification		II.1.2.	-		ards responsibilities, operation Annex I to Commission Deleg	-
II: Certi	(1) ° [II.1.	animals de	scribed in Par		ryos] (1) 🛘 [micromanipulated or produced, processed and	
art		II.1.1.	is approved a	and kept in a register l	by the competent authority;	
P		II.1.2.			ards responsibilities, operationd 3 of Annex I to Delegated R	
	II.2.		es] (1) [embry com donor ani		rt I are intended for artificial	reproduction and were
		II.2.1.			ee birth in the Union, or have for entry into the Union;	entered the Union in
		(1)(3) □ [II.2.2.	disease virus		e thereof which is free from ir d eradication programme for	
		II.2.3.			mber State or zone thereof, or uthority in a third country or	
			II.2.3.1.	animals has not been collection of the \Box [or	th Brucella abortus, B. melited reported during the last 42 do locytes] (1) [embryos] (1), a e date of collection of the [] [ays prior to the date of and in which during at least
			(1) □ either		and risk mitigating measure of Commission Delegated Regduced;	
			(1) □ and/or	suis has be establishm	ce for infection with Brucella een carried out on the porcine ents in accordance with Artic Regulation (EU) 2020/688;]	animals kept in the
			II.2.3.2.	with Aujeszky's disea	ological, virological or pathol se virus had been detected du ollection of the [] [oocytes] (1]	uring at least 12 months
		II.2.4.		of transmissible anim	narian or a team member and nal diseases on the day of colle	
		II.2.5.		d as provided for in Ar EU) 2019/2035;	rticle 52 or Article 54(2) of Cor	nmission Delegated
		II.2.6.		O days prior to the dat he collection period;	e of first collection of the \Box [o	oocytes] (1) 🗆 [embryos] (1)
			II.2.6.1.	restrictions established infection with rinder	nments situated in a zone not ed due to the occurrence of fo pest virus, classical swine fevo se relevant for porcine anima	ot and mouth disease, er or African swine fever, or
			II.2.6.2.	melitensis and B. suis Aujeszky's disease vii	establishment where infections, infections with rabies virus, a rus and infection with porcine e virus have not been reported	anthrax, infection with e reproductive and

en 3/6

	NIOIN				2021/10	44 (2021/403) N		CCCTIES	
	II. Health informat	10n							
			II.2.6.3.	subject to in point II	movement r	th animals from estrictions due to n establishments 2.6.2;	o the occurr	ence of disea	ises referred to
п			II.2.6.4.	were not	used for natu	ıral breeding;			
ıtio	II.2	2.7.	comply with	the follow	ing conditior	ns as regards foo	t and mouth	disease:	
fice			II.2.7.1.	they come	e from establ	ishments			
Part II: Certification			-	a 10-km r	adius centre	ere foot and mou d on the establish lection of the □	nment for at	least 30 day	-
Part			-		nmediately p	th disease has no rior to the date o	_	_	
	(1) eitl	o her	[II.2.7.2.	they were	not vaccina	ted against foot a	and mouth d	isease;]	
	(1)	(4) or	[II.2.7.2.	-		against foot and i		_	months prior
				II.2.7.2.1.		en vaccinated ag s immediately pı			
				II.2.7.2.2.	that complied 1(b), of Ann complies wi	used for fertilisat es with the condi ex II to Delegated th the conditions Delegated Regula	tions set out d Regulation s set out in P	t in Part 5, Cl (EU) 2020/6 Part 5, Chapte	napter I, point 86 or the semen
				II.2.7.2.3.	-	ezing, the embryoried out in accor cl (5);		•	
				II.2.7.2.4.	date of colle	s were stored de ection, and durin cal signs of foot a	g this period	d the donor a	days from the inimals have not
		2.8.	syndrome vi	rus, with n	egative resu	for infection with lts, on two occasi ed within 15 day	ions, at an ir	nterval of no	t less than 21
	II.3. The	e 🗆 [ood	ytes] (1) □ [embryos]	(1) described	in Part I:			
	II.3			[Part 2] (1)	□ [Part 3] (stored in accorda 1) □ [Part 4] (1)			
	II.3			s provided	for in Articl	ges on which the e 10 of Delegated			
	II.3	3.3.	are transpor	ted in a co	ntainer whic	h:			
			II.3.3.1.	collection	or production	ered prior to the on team under re rian, and the sea	sponsibility	of the team	veterinarian, or
			II.3.3.2.	has been container		either disinfected	d or sterilise	d before use	, or is single-use
			(1)(7)II.3.3.3. □ [has been if		a cryogenic agei	nt which has	s not been pr	eviously used
	(1)	(8) 🗆	are placed in	straws or	other packag	ges which are sec	curely and h	ermetically	sealed;

en 4/6

	II. Health info	rmation			
		[II.3.4.			
		II.3.5.	are transported in a container whe physical compartments or by being		
Part II: Certification	(1)(9) □ [II.4.	embryos] (semen colle approved f Member St Annex XI to and stored	vivo derived embryos] (1) [in vit. 1) described in Part I were conceive ection centre, germinal product proor the collection, processing and/or ate or by the competent authority of Commission Implementing Regula in accordance with the requirement ated Regulation (EU) 2020/686.]	ed by artificial insemination uncessing establishment or gerications of semen by the complete fathird country or territory, ation (EU) 2021/404, and which	ssing semen coming from a minal product storage centre petent authority of a or zone thereof listed in h was collected, processed
Part	(1)(10) □ [II.5.		ng antibiotic or mixture of antibiot storage media:	ics (11) has been added to the	collection, processing,
	II.6.	The □ [oo	cytes] (1) 🛘 [embryos] (1) described	l in Part I are dispatched:	
	(1) either	movement for those sp restrictions were estab	l [embryo collection team] (1) [er restrictions affecting porcine animal pecies or diseases subject to emerge ado not apply to these oocytes or en lished, and they have not been in contaquate period.]	als and established for reasor ncy measures relevant for th nbryos because they were col	ns of listed diseases relevant ose species, or those lected before the restrictions
	(1) or	movement	l [embryo collection team] (1)	als and established for	r from a zone subject to (12), but derogations
	(1)	☐ [they con	mply with the requirements set out	in (13);]]	
	(1)	\square [and in p	particular, they are (14	4).]]	
	from the E Protocol or animal hea This anima	uropean Un n Ireland/No alth certifica al health cer	Agreement on the withdrawal of the Agreement on the withdrawal of the ion and the European Atomic Energy rthern Ireland in conjunction with the include the United Kingdom in retificate shall be completed in according to fannex I to Commission Imple	gy Community, and in particu Annex 2 to that Protocol, refe espect of Northern Ireland. Iance with the notes for the c	lar Article 5(4) of the rences to the Union in this ompletion of certificates
	Part I:				
	Box reference I.11:		spatch": Indicate the unique appro or production team of dispatch of th		
	Box reference I.12:		estination": Indicate the address and ent of destination of the consignment		oval number of the
	Box reference I.19:	Seal numb	er shall be indicated.		
	Box reference I.26:	Total numl	per of packages shall correspond to	the number of containers.	
	Box reference I.30:		cify if in vivo derived embryos, in v pulated embryos.	vivo derived oocytes, in vitro	produced embryos or
			"Identification number": Indicate i	dentification number of each	donor animal.
			"Identification mark": Indicate ma embryos of the consignment are pl	_	ages where the oocytes or

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Consignment we "Approval or re approval numb the oocytes or e "Quantity": Indipersion of the consignment of the Council and Articles and of the Council and Articles and of the International for the use of embryo trans International Embryo Techn (http://www.iets.org/). (6) Applicable for in vivo derive (Applicable for consignment micromanipulated embryos (9) Does not apply to oocytes. (10) Mandatory attestation in case (Approval or reasonable) (10)	•
Consignment we "Approval or re approval numb the oocytes or e "Quantity": Individual Part II: (1) Delete if not applicable. (2) Only embryo collection or pregister referred to in Articiand of the Council and Articiand of the Council and Articiand of the International for the use of embryo trans International Embryo Techn (http://www.iets.org/). (6) Applicable for in vivo deriv (Applicable for consignment micromanipulated embryos (9) Does not apply to oocytes. (10) Mandatory attestation in care	rere collected or produced. registration number of plant/establishment/centre": Indicate the unique our of the embryo collection team or the embryo production team by which embryos of the consignment were collected or produced. Iticate number of straws or other packages with the same mark. production teams approved by the competent authority and included in the cle 101(1), point (b), of Regulation (EU) 2016/429 of the European Parliament icle 7 of Delegated Regulation (EU) 2020/686. Iderived embryos subject to trypsin treatment. The consignment of in vivo derived embryos. The Embryo Technology Society — A procedural guide and general information after technology emphasising sanitary procedures, published by the molgy Society, 1 111 North Dunlap Avenue, Savoy, Illinois 61 874, USA aved embryos. The Embryos. The Embryos. The Embryos.
"Approval or re approval numb the oocytes or e "Quantity": Indi Part II: (1) Delete if not applicable. (2) Only embryo collection or pregister referred to in Articiand of the Council and Articiand of the Council and Articiand of the International for the use of embryo trans International Embryo Technicational Embryo Technication In Capplicable for consignment micromanipulated embryos Technication In Capplication In Capplicat	egistration number of plant/establishment/centre": Indicate the unique per of the embryo collection team or the embryo production team by which embryos of the consignment were collected or produced. Iticate number of straws or other packages with the same mark. production teams approved by the competent authority and included in the cle 101(1), point (b), of Regulation (EU) 2016/429 of the European Parliament icle 7 of Delegated Regulation (EU) 2020/686. Iderived embryos subject to trypsin treatment. The consignment of in vivo derived embryos. The Embryo Technology Society — A procedural guide and general information after technology emphasising sanitary procedures, published by the molgy Society, 1 111 North Dunlap Avenue, Savoy, Illinois 61 874, USA wed embryos. The Embryos. The Embryos. The Embryos.
approval numb the oocytes or e "Quantity": Indi Part II: (1) Delete if not applicable. (2) Only embryo collection or p register referred to in Artic and of the Council and Artic and of the Council and Artic (3) Not applicable for in vivo d (4) Option available only for th (5) Manual of the International for the use of embryo trans International Embryo Tech (http://www.iets.org/). (6) Applicable for in vivo deriv (7) Applicable for frozen oocyt (8) Applicable for consignment micromanipulated embryos (9) Does not apply to oocytes. (10) Mandatory attestation in ca	per of the embryo collection team or the embryo production team by which embryos of the consignment were collected or produced. Idicate number of straws or other packages with the same mark. Production teams approved by the competent authority and included in the cle 101(1), point (b), of Regulation (EU) 2016/429 of the European Parliament icle 7 of Delegated Regulation (EU) 2020/686. Iderived embryos subject to trypsin treatment. The consignment of in vivo derived embryos. Final Embryo Technolgy Society — A procedural guide and general information after technology emphasising sanitary procedures, published by the sanolgy Society, 1 111 North Dunlap Avenue, Savoy, Illinois 61 874, USA ved embryos. The consignment of the
(4) Option available only for the use of embryo trans International Embryo Technical Embryo	production teams approved by the competent authority and included in the cle 101(1), point (b), of Regulation (EU) 2016/429 of the European Parliament icle 7 of Delegated Regulation (EU) 2020/686. derived embryos subject to trypsin treatment. he consignment of in vivo derived embryos. al Embryo Technolgy Society — A procedural guide and general information sfer technology emphasising sanitary procedures, published by the anolgy Society, 1 111 North Dunlap Avenue, Savoy, Illinois 61 874, USA ved embryos. tes or embryos.
(4) Option available only for the use of embryo trans International Embryo Technical Embryo	cle 101(1), point (b), of Regulation (EU) 2016/429 of the European Parliament icle 7 of Delegated Regulation (EU) 2020/686. derived embryos subject to trypsin treatment. he consignment of in vivo derived embryos. al Embryo Technolgy Society — A procedural guide and general information sfer technology emphasising sanitary procedures, published by the anolgy Society, 1 111 North Dunlap Avenue, Savoy, Illinois 61 874, USA ved embryos. tes or embryos.
(4) Option available only for the use of embryo trans International Embryo Technical Embryo	cle 101(1), point (b), of Regulation (EU) 2016/429 of the European Parliament icle 7 of Delegated Regulation (EU) 2020/686. derived embryos subject to trypsin treatment. he consignment of in vivo derived embryos. al Embryo Technolgy Society — A procedural guide and general information sfer technology emphasising sanitary procedures, published by the anolgy Society, 1 111 North Dunlap Avenue, Savoy, Illinois 61 874, USA ved embryos. tes or embryos.
(4) Option available only for the use of embryo trans International Embryo Technical Embryo	cle 101(1), point (b), of Regulation (EU) 2016/429 of the European Parliament icle 7 of Delegated Regulation (EU) 2020/686. derived embryos subject to trypsin treatment. he consignment of in vivo derived embryos. al Embryo Technolgy Society — A procedural guide and general information sfer technology emphasising sanitary procedures, published by the anolgy Society, 1 111 North Dunlap Avenue, Savoy, Illinois 61 874, USA ved embryos. tes or embryos.
(4) Option available only for the use of embryo trans International Embryo Technical Embryo	he consignment of in vivo derived embryos. al Embryo Technolgy Society — A procedural guide and general information sfer technology emphasising sanitary procedures, published by the nnolgy Society, 1 111 North Dunlap Avenue, Savoy, Illinois 61 874, USA ved embryos. tes or embryos.
(4) Option available only for the (5) Manual of the International for the use of embryo trans International Embryo Techn (http://www.iets.org/). (6) Applicable for in vivo deriv (7) Applicable for frozen oocytes) Applicable for consignment micromanipulated embryos (9) Does not apply to oocytes. (10) Mandatory attestation in care	he consignment of in vivo derived embryos. al Embryo Technolgy Society — A procedural guide and general information sfer technology emphasising sanitary procedures, published by the nnolgy Society, 1 111 North Dunlap Avenue, Savoy, Illinois 61 874, USA ved embryos. tes or embryos.
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for the use of embryo trans International Embryo Tech (http://www.iets.org/). (6) Applicable for in vivo deriv (7) Applicable for frozen oocyt (8) Applicable for consignment micromanipulated embryos (9) Does not apply to oocytes. (10) Mandatory attestation in ca	sfer technology emphasising sanitary procedures, published by the nnolgy Society, 1 111 North Dunlap Avenue, Savoy, Illinois 61 874, USA ved embryos. tes or embryos.
 (7) Applicable for frozen oocyte (8) Applicable for consignment micromanipulated embryos (9) Does not apply to oocytes. (10) Mandatory attestation in ca 	tes or embryos.
 (8) Applicable for consignment micromanipulated embryos (9) Does not apply to oocytes. (10) Mandatory attestation in ca 	•
micromanipulated embryos (9) Does not apply to oocytes. (10) Mandatory attestation in ca	its where oocytes, in vivo derived embryos, in vitro produced embryos and
(10) Mandatory attestation in ca	os of porcine animals are placed and transported in one container.
,	
(11) In court the manager of the cour	ase antibiotic(s) was/were added.
(11) Insert the name(s) of the an	ntibiotic(s) added and its (their) concentration.
(12)Insert the name of the disease(s).	
(13) Insert the specific reference Commission providing for t	ce to the article(s), title, and number of the relevant legal act(s) adopted by the those requirements.
	on(s) provided for in and required by the relevant legal act(s) adopted by the in Article 159(2), points (a), (b) and (c), of Regulation (EU) 2016/429.
Certifying Officer/Official veterinarian	
Name (in capital letters) Date of declaration	Qualification and title Signature
Stamp	Signature

6 / 6