								1111111	
	I.1. Consignor				I.2. IMSOC ref	erence	I.2.a. Local reference		
	Name				I.3. Central Competent Autho			nt Authority	
	Address				I.4. Local Competent Authori			Authority	
	Country		ISO Code						
-	1.5. 0				10.0	I.6. Operator conducting assembly operations independently of an			
님	I.5. Consignee				establishmen	conducting assembly of	perations independen	tly of an	
Part I: Description of consignment	Name								
2	Address				Name Address				
50	Country		ISO Code		Approval Nu:	mhor			
ns					Country	linder	ISO Code		
ខ					country		150 Coue		
Ч	I.7. Country of orig	gin		ISO Code	I.9. Country of	f destination		ISO Code	
g									
E I					140 D				
음	I.8. Region of origi			Code	I.10. Region of			Code	
S	I.11. Place of dispatch I.					lestination			
ĕ	Name				Name				
$\overline{\cdot}$	Address				Address				
ť	Approval Number	r			Approval Number				
Pai	Country		ISO Code		Country		ISO Code		
ŀ	I 12 Dloop of last	nď			114 Data 3	time of department			
	I.13. Place of loadi	пg			1.14. Date and	time of departure			
	Name					<b>A</b>			
	Address								
	Approval Number	r							
	Country		ISO Code						
-	I.15. Means of Trai	nsnort			I.16. Transpor	ter			
		1	T. 1						
	transport document				Name				
					Address				
					Approval Nu	mber			
					Country		ISO Code		
					L17. Accompa	nying documents			
					[en] accompanyi				
					accompanyi ng.documen t.document.		Date of issue		
					number				
							Place of		
-					Country		issue		
	I.18. Transport cor	nditions		_	· · · · □				
	Chilled 📙		Frozen L		Ambient 📙				
-	140.0	(C 1) .							
	I.19. Container No	/ Seal No							
ŀ	I.20. Certified as								
	Germinal products	. П							
	Germinal producta	, <b>ш</b>							
ļ	I.21. For transit th	rough a third cour	ntry						
	Third country	-			ISO Code				
	Exit point				BCP code				
	Entry point				BCP code				
	I.22. For transit th	rough Member Sta	ite(s)		I.23. For expo	rt			
	Member State	0	ISO Code		Third country		ISO Code		
					Exit point BCP code				
	I.24. Estimated jou	rney time			I.25. Journey	Log			
╞									
						intity			
ŀ	I.30. Description of	f consignment		1					
		-	26	Idoptification	Number	Quantity	Noture of com	amodity	
	Commodity	Specie	28	Identification	numper	Quantity	Nature of cor	innoany	
			1				I		
	Identification Mar	rk	Package count		Date of collect	tion / production	Plant / Establishmen	t / Centre	
ſ									

						20.	1/403 DO		
	II. Health information								
	I, the undersigned official veterinarian, hereby certify that:								
	(1)	□ [II.1.	[II.1. The in vivo derived embryos of bovine animals described in Part I have been collected, processed and stored, and dispatched by the embryo collection team(2) which						
			II.1.1.	is approve	n a register by the o	competent	authority;		
cation			II.1.2.	-	ind equipmer	nents as regards responsibilities, operational procedures, nt set out in Part 2 of Annex I to Delegated Regulation (EU)			
Part II: Certification	(1)	🗆 [II.1.	The oocytes(1)/ in vitro produced embryos(1)/ micromanipulated embryos(1) of bovine animals described in Part I have been collected or produced, processed and stored, and dispatched by the embryo production team(2) which						
art ]			II.1.1.	is approve	ed and kept in	n a register by the o	competent	authority;	
P			II.1.2.	facilities a	-	nents as regards re nt set out in Parts 2 86.]	-		•
		II.2.	The oocytes(1)/ embryos(1) described in Part I are intended for artificial repro- were obtained from the donor animals which						ction and
			II.2.1.			mained since birth ith the requiremer			
			II.2.2.	establishn	nents under o	ents in a Member S official control by th a zone thereof			
				II.2.2.1.	bovis, M. ca	nfection with Myco aprae and M. tuber in any establishme	culosis), an	d they have nev	er been kept
				II.2.2.2.		nfection with Bruce ave never been kep h status;			
	(1)		∘ either	[II.2.2.3.		nzootic bovine leul in any establishme			-
	(1)		° or	[11.2.2.3.	responsible has been ne	om enzootic bovine e for the establishm o clinical case of en he preceding 3 yea	ent of orig	in has certified t	hat there
	(1)		○ either	[II.2.2.4.	vulvovagin	nfectious bovine rh itis and they have a ent of a lower healt	never been	-	
	(1)		o or	[II.2.2.4.	vulvovagin establishm case of infe	m infectious bovin itis and the official ent of origin has ce ctious bovine rhin itis during a period	veterinari rtified that otracheitis/	an responsible fo there has been t infectious pustu	or the no clinical lar
				II.2.2.5.	the 30 day	ırra (Trypanosoma period prior to coll embryos(1), and			
	(1)			○ either		not been reported i to collection(1)/ pr ;]			
	(1)			∘ or	years prior embryos(1)	been reported in th to collection(1)/ pr and following the under movement re	oduction(1 last outbre	) of the oocytes(2 eak the establishi	1)/
								als have been re hment, and	emoved

	II. Health information						
Part II: Certification					-	have been subject (Trypanosoma evidiagnostic method Annex I to Comm (EU) 2020/688, caresults, on sampling after the infected	timals on the establishment eted to a test for surra vansi) with one of the ods provided for in Part 3 of hission Delegated Regulation rried out, with negative es taken at least 6 months animals have been e establishment;]
rt II: Cer		II.2.3.	symptoms	or clinical si	igns of tran		member and did not show liseases on the day of os(1);
Pai		II.2.4.		ually identi (EU) 2019/2	-	vided for in Article	38 of Commission Delegated
		II.2.5.	-			or to the date of fir mbryos(1) and dur	rst collection(1)/ ing the collection period
			II.2.5.1.	established infection w contagious	l due to the vith rinderr bovine ple	occurrence of foot best virus, infectior	d in a restricted zone t-and-mouth disease, n with Rift Valley fever virus, lumpy skin disease or of an caprine animals;
			II.2.5.2.	abortus, B. tuberculosi rabies, anti leukosis, in vulvovagin haemorrha (serotypes	melitensis is complex hrax, surra ifectious bo itis, bovine igic disease 1-24) have	and B. suis, infecti (M. bovis, M. capra (Trypanosoma eva ovine rhinotracheit e viral diarrhoea, in e virus and infectio not been reported;	
			II.2.5.3.	restricted z	zone due to from establ	the occurrence of ishments which do	stablishments situated in a diseases referred to in point o not meet the conditions
			II.2.5.4.	were not u	sed for nat	ural breeding;	
		II.2.6.	comply wit	th the follow	ving conditi	ions as regards foo	t-and-mouth disease
			II.2.6.1.	they come	from estab	lishments	
					-	disease has not b radius centred or period of at least	ea where foot-and-mouth een reported within a 10-km n the establishment for a 30 days immediately prior lection of the oocytes(1)/
					-	been reported du months immedia	d-mouth disease has not uring a period of at least 3 tely prior to the date of pocytes(1)/ embryos(1);
	(1)	$\circ$ either	[II.2.6.2.	they were a	not vaccina	ited against foot-an	id-mouth disease;]
	(1)(3) • or [II.2.6.2.			they were vaccinated against foot-and-mouth disease during the period of 12 months prior to the date of collection or production of the embryos and			
				II.2.6.2.1.	within the		ainst foot-and-mouth disease 30 days immediately prior to embryos;

II. Health information				
			II.2.6.2.2.	the semen used for fertilisation was collected from a male donor that complies with the conditions set out in point 1(b) or the semen complies with the conditions set out in point 2 of Chapter I of Part 5 of Annex II to Delegated Regulation (EU) 2020/686;
			II.2.6.2.3.	prior to freezing, the embryos have been subjected to trypsin washing carried out in accordance with the recommendations of the IETS Manual(4);
			II.2.6.2.4.	the embryos were stored deep frozen for a period of at least 30 days from the date of collection, and during this period the donor animal has not shown clinical signs of foot-and-mouth disease;]
(1)(5)	□ [II.2.7.		th at least or e virus (sero	ne of the following conditions as regards infection with types 1-24):
(1)	□ either	[II.2.7.1.	during coll thereof fre where no c	been kept for a period of at least 60 days prior to and ection of the oocytes in a third country, territory or zone e from infection with bluetongue virus (serotypes 1-24) case of infection with bluetongue virus (serotypes 1-24) has rmed during the last 24 months in the targeted animal ;]
(1)	□ and/or	[II.2.7.2.	seasonally to and duri or zone the	been kept in a seasonally disease-free zone, during the disease-free period, for a period of at least 60 days prior ing collection of the oocytes, in a third country, territory ereof with an approved eradication programme against with bluetongue virus (serotypes 1-24);]
(1)	□ and/or	[II.2.7.3.	seasonally to and duri or zone the of the cons obtained th Member St that season	been kept in a seasonally disease-free zone, during the disease-free period, for a period of at least 60 days prior ing collection of the oocytes, in a third country, territory ereof where the competent authority of the place of origin ignment of oocytes(1)/ in vitro produced embryos(1) has he prior written consent of the competent authority of the ate of destination to the conditions for establishment of hally disease-free zone and to accept the consignment of in vitro produced embryos(1);]
(1)	□ and/or	[II.2.7.4.	-	been kept in a vector-protected establishment for a period 50 days prior to and during collection of the oocytes;]
(1)	□ and/or	[II.2.7.5.	the bluetor	been subjected to a serological test to detect antibodies to ngue virus serogroup 1-24, with negative results, between lays from the date of each collection of the oocytes;]
(1)	□ and/or	[II.2.7.6.	bluetongue	been subjected to an agent identification test for e virus (serotypes 1-24), with negative results, on blood en on the day of collection of the oocytes;] ]
(1)(5)	□ [II.2.8.			ne of the following conditions as regards infection with c disease virus (serotypes 1-7) (EHDV 1-7):
(1)	□ either	[II.2.8.1.	during coll thereof wh	been kept for a period of at least 60 days prior to and ection of the oocytes in a third country, territory or zone ere EHDV 1-7 has not been reported for a period of at receding 2 years within a radius of 150 km of the ent;]
(1)	□ and/or	[II.2.8.2.	-	been kept in a vector-protected establishment for a period 50 days prior to and during collection of the oocytes;]
(1)	□ and/or	[II.2.8.3.	findings th have been	ent in the exporting country in which according to official e following serotypes of EHDV exist: and subjected with negative results in each case to the ests carried out in an official laboratory:

II. Health information							
		□ either	[II.2.8.3.1.	negative results, on blood say	mple taken between 28 and		
		□ and/or	[II.2.8.3.2.				
	□ [II.2.9.			-	in Chapter III of Part 1 of		
II.3.	The oocyte	es(1)/ embry	os(1) descrik	oed in Part I			
	II.3.1.	requireme	nas been collected, processed and stored in accordance with animal health requirements set out in Part 2(1)/Part 3(1)/Part 4(1)/Part 5(1) and Part 6 of Annex III to Delegated Regulation (EU) 2020/686;				
	II.3.2.	accordanc	le 10 of Delegated Regulation				
	II.3.3.	are transp	orted in a co	ontainer which:			
		II.3.3.1.	collection o veterinaria	or production team under resp an, or by an official veterinaria	oonsibility of the team		
		II.3.3.2.			or sterilised before use, or is		
		□ [II.3.3.3.	has been fi	lled in with the cryogenic age	nt which not have been		
	□ [II.3.4.	are placed sealed;		-	curely and hermetically		
	II.3.5.			· · ·	-		
□ [[II.4.	embryos(1 coming fro germinal p semen by third coun	in vivo derived embryos(1)/ in vitro produced embryos(1)/ micromanipuryos(1) described in Part I were conceived by artificial insemination using from a semen collection centre, germinal product processing establishinal product storage centre approved for the collection, processing and, en by the competent authority of a Member State or by the competent authority of a Member State or by the competent authority of a Member State or by the competent authority of a Member State or by the competent authority of a Member State or by the competent authority of a Member State or by the competent authority of a Member State or by the competent authority of a Member State or by the competent authority of a Member State or by the competent authority of a Member State or by the competent authority of a Member State or by the competent authority of a Member State or by the competent authority of a Member State or by the competent authority of a Member State or by the competent authority of a Member State or by the competent authority of a Member State or by the competent authority of a Member State or by the competent authority of a Member State or by the competent authority of a Member State or by the competent authority of a Member State or by the competent authority of a Member State or by the competent authority of a Member State or by the competent authority of a Member State or by the competent authority of a Member State or by the competent authority of a Member State or by the competent authority of a Member State or by the competent authority of a Member State or by the competent authority of a Member State or by the competent authority of a Member State or by the competent authority or a Member State or by the competent authority of a Member State or by the competent authority or a Member State or by the competent authority or a Member State or by the competent authority or a Member State or by the competent authority or a Member State or by the competent authority or a Member State or by the competent authority o					
□ [II.5.					dded to the collection,		
	II.3.	□ [II.2.9. II.3. The oocyte II.3.1. II.3.2. II.3.2. II.3.3. □ [II.3.4. II.3.5. □ [[II.4. The in vive embryos(1) coming fro germinal p semen by third coun Regulation □ [II.5. The follow	□ either     □ and/or     □ [II.2.9.   comply with Annex II to Annex II to Delege     II.3.   The oocytes(1)/ embry     II.3.   The oocytes(1)/ embry     II.3.1.   has been of requirements     II.3.2.   are placed accordance     II.3.2.   are placed accordance     II.3.3.   are transp     II.3.3.   II.3.3.1.     II.3.3.   II.3.3.2.     □ [II.3.4.   are placed sealed;     II.3.5.   are transp     physical co   sealed;     II.3.5.   are transp  <	□ either   [II.2.8.3.1.     □ and/or   [II.2.8.3.2.     □ [II.2.9.   comply with animal he Annex II to Delegated I     II.3.   The oocytes(1)/ embryos(1) descrift     II.3.   The oocytes(1)/ embryos(1) descrift     II.3.   has been collected, progrequirements set out in III to Delegated Regula     II.3.   are placed in straws or accordance with require(EU) 2020/686 and that     II.3.3.   are transported in a concollection of veterinaria number as     II.3.3.   are transported in a concollection of veterinaria number as     II.3.3.   has been collection of veterinaria number as     II.3.3.   are placed in straws or sealed;     II.3.3.   has been fill     II.3.3.   has been fill     II.3.3.   has been fill     II.3.5.   are transported in a consealed;     II.3.6.   are transported in a consealed;     II.3.7.   The in vivo derived embryos(1) in embryos(1) described in Part I web coming from a semen collection consealed;     II.1.4.   The in vivo derived embryos(1) in embryos(1) described in	either   [II.2.8.3.1.]   a serological test to detect an negative results, on blood sam 60 days from the date of the eight results, on blood sam 60 days from the date of the eight results, on blood sample take the oocytes.]]]     II.2.9.   comply with animal health requirements laid down Annex II to Delegated Regulation (EU) 2020/686.]     II.3.   The oocytes(1)/ embryos(1) described in Part I     II.3.   has been collected, processed and stored in accordar requirements set out in Part 2(1)/Part 3(1)/Part 4(1)/F III to Delegated Regulation (EU) 2020/686;     II.3.   are placed in straws or other packages on which the accordance with requirements provided for in Articl (EU) 2020/686 and that mark is indicated in Box I.30;     II.3.3.   are transported in a container which:     II.3.3.1.   was sealed and numbered prior to the di collection or production team under resp veterinarian, or by an official veterinaria number as indicated in Box I.19;     II.3.3.2.   has been cleaned and either disinfected or single-use container;     II.3.3.1.   was sealed and numbered prior to the di collection or production team under resp veterinarian, or by an official veterinaria number as indicated in Box I.19;     II.3.3.   has been cleaned and either disinfected or single-use container;     II.3.3.   has been filled in with the cryogenic age previously used for other products.]     II.3.5.   are transported in a container where they are separiphysical compartments or by being placed in second compry, second		

	II. Health info	rmation								
Part II: Certification	Notes	Notes								
	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:									
	Box reference I.11:	"Place of dispatch": Indicate the unique approval number and the name and address of the embryo collection or production team of dispatch of the consignment of oocytes or embryos.								
	Box reference I.12:									
	Box reference I.19:	Seal number shall be indicated.								
	Box reference I.26:	Total number of packages shall correspond to the number of containers.								
Box "Species": select amongst "Bos taurus", "Bison bison" or "Bubalus bubalis" as appropriate. reference I.30:										
		"Type": specify if oocytes, in vivo derived embrees.	ryos, in vitro produced embr	yos or micromanipulated						
		"Species": select amongst "Ovis aries" or "Capra hircus" as appropriate.								
	"Identification number": Indicate identification number of each donor animal.									
		"Identification mark": indicate mark on the st consignment are placed.	raw or other packages where	oocytes or embryos of the						
		"Date of collection/production": indicate the d collected or produced.	ate on which oocytes or emb	ryos of the consignment was						
		"Approval or registration number of plant/est of the embryo collection or production team by produced.	y which the oocytes or embry	ros were collected or						
		"Quantity": Indicate number of straws or othe	r packages with the same ma	rk.						
	Part II:									
	(1)	Delete if not applicable.								
	(2)	Only embryo collection or production teams approved by the competent authority and included in the register referred to in Article 101(1)(b) of Regulation (EU) 2016/429 and Article 7 of Delegated Regulation (EU) 2020/686.								
	(3)	Option available only for the consignment of in vivo derived embryos.								
	(4)	Manual of the International Embryo Transfer Society — A procedural guide and general information for the use of embryo transfer technology emphasising sanitary procedures, published by the International Embryo Transfer Society, 1 111 North Dunlap Avenue, Savoy, Illinois 61 874 USA (http://www.iets.org/).								
	(5)	Applicable for the consignment of oocytes and	in vitro produced embryos.							
	(6)	Applicable for frozen oocytes or embryos.								
	(7)	Applicable for the consignment where in one of produced embryos and micromanipulated embryos and microm	-	-						
	(8)	Does not apply to oocytes.								
	(9)	Mandatory attestation in case antibiotics were added.								
	(10)	Insert the name(s) of the antibiotic(s) added ar	nd its(their) concentration.							
Certifying Officer/Official veterinarian										

## 2021/403 BOV-OOCYTES-EMB-A-INTRA

	II. Health information		
	Name (in capital letters) Date of signature Stamp	Qualification and title Signature	
Part II: Certification	Date of signature Stamp	Qualification and title Signature	