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 Transport				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 document			
					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						
	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						
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	II. Health info	rmation									
	I, the undersigned official veterinarian, hereby certify that:										
	II.1.	The germinal product storage centre (1) described in box I.11 at which the \square [semen] (2) \square [oocytes] (2) \square [in vivo derived embryos] (2) \square [in vitro produced embryos] (2) \square [micromanipulated embryos] (2) was/were stored:									
ц		II.1.1.	is approve	d and kept in a register l	by the competent authority;						
rtificatio	II.2. (2) o either	II.1.2.	complies with requirements as regards responsibilities, operational procedures, facilities and equipment set out in Part 5 of Annex I to Commission Delegated Regulation (EU) 2020/686;								
II: Ce	II.2.	The \square [semen] (2) \square [oocytes] (2) \square [in vivo derived embryos] (2) \square [in vitro produced embryos] (2) \square [micromanipulated embryos] (2) described in Part I is/are dispatched from:									
Part	(2) o either	restrictions species or o not apply t established produced of	s affecting b diseases sub o these gerr l, and the [2 embryos] (2)	d in box I.11 or a zone not sub plished for reasons of listed d ures relevant for those specie they were collected before the [s] (2) [[in vivo derived embed] embryos] (2) has/have not be an adequate period.]	iseases relevant for those es, or those restrictions do e restrictions were bryos] (2) [in vitro]						
	(2) ∘ or	affecting b	the germinal product storage centre described in box I.11 or a zone subject to movement restrictions affecting bovine animals and established for (3), but derogations from movement restrictions have been granted, and:								
	(2)	□ [it/they o	comply(ies)	with the requirements s	et out in (4);]]						
	(2)	□ [and in	☐ [and in particular, it/they is/are (5).]]								
II.3. The □ [semen] (2) □ [oocytes] (2) □ [in vivo derived embryos] (2) □ [in vitro produced en □ [micromanipulated embryos] (2) described in Part I is/are intended for artificial reproduced en □ [micromanipulated embryos] (2) described in Part I is/are intended for artificial reproduced en □ [micromanipulated embryos] (2) described in Part I is/are intended for artificial reproduced en □ [micromanipulated embryos] (2) □ [in vivo derived embryos] (3) □ [in vitro produced en □ [micromanipulated embryos] (4) □ [in vivo derived embryos] (5) □ [in vitro produced en □ [micromanipulated embryos] (6) □ [in vivo derived embryos] (7) □ [in vitro produced en □ [micromanipulated embryos] (8) □ [in vivo derived embryos] (9) □ [in vitro produced en □ [micromanipulated embryos] (9) □ [in vivo derived											
	(2) ☐ [II.3.1. has/have been ☐ [collected] (2) ☐ [produced] (2), ☐ [processed] (2) ☐ [stored] (2) ☐ [ir semen collection centre] (2) (6) ☐ [by an embryo collection team] (2) (6) ☐ [by an embryo production team (2) (6) ☐ [and] (2) ☐ [processed] (2) ☐ [stored] (2) ☐ [in a germinal product processing establishment] (2) (6) ☐ [and stored in a germinal product storage centre] (2) (6) situated in the Member State of its/their collection or production and complying with requirements as regards responsibilities, operational procedures, facilities and equipment set out in ☐ [Part 1] (2) ☐ [Part 2] (2) ☐ [Part 3] (2) ☐ [Part 4] (2) ☐ [Part 5] (2) of Annex I to Delegated Regulation (EU) 2020/686, and was/were moved to the germinal product storage centre indicated in box I.11 situated in the Member State of its/their collection or production under animal health certification requirements at least a strict as those provided for in:										
	(2)		□ either	[Model BOV-SEM-A-IN]	ΓRA (7);]						
	(2)		□ and/or	[Model BOV-SEM-B-IN]	TRA (7);]						
	(2)		□ and/or	[Model BOV-SEM-C-IN]	TRA (7);]						
	(2)		□ and/or	[Model in Annex D1 to	Directive 88/407/EEC(7);]						
	(2)		\square and/or	[Model in Annex D2 to	Directive 88/407/EEC(7);]						
	(2)		\square and/or	[Model in Annex D3 to	Directive 88/407/EEC(7);]						
	(2)		\square and/or	[Model BOV-OOCYTES-	EMB-A-INTRA(7);]						
	(2)		\square and/or	[Model BOV-EMB-B-IN]	ΓRA(7);]						
	(2)		\square and/or	[Model BOV-GP-PROCE	SSING-INTRA(7);]						
	(2)		\square and/or	[Model BOV-GP-STORA	GE-INTRA(7);]]						
	(2) □ and/or	[II.3.1. has/have been \Box [collected] (2) \Box [produced] (2) \Box [processed] (2) \Box [stored] (2) \Box [in a									

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	II. Health info	rmation				
			51 (0)		1	
			germinal p	product storage centre ir	lation (EU) 2020/686, and was adicated in box I.11 situated in tificate(s) in accordance with	n another Member State
	(2)		\square either	[Model BOV-SEM-A-IN]	ΓRA (7);]	
ď	(2)		\square and/or	[Model BOV-SEM-B-IN]	TRA (7);]	
atic	(2)		\square and/or	[Model BOV-SEM-C-IN]	TRA (7);]	
ific	(2)		□ and/or	[Model in Annex D1 to	Directive 88/407/EEC (7);]	
ert	(2)		\square and/or	[Model in Annex D2 to	Directive 88/407/EEC (7);]	
Part II: Certification	(2)		\square and/or	[Model in Annex D3 to	Directive 88/407/EEC (7);]	
rt I	(2)		\square and/or	[Model BOV-OOCYTES-	EMB-A-INTRA (7);]	
Pa	(2)		\square and/or	[Model BOV-EMB-B-IN	ΓRA (7);]	
	(2)		\square and/or	[Model BOV-GP-PROCE	SSING-INTRA (7);]	
	(2)		\square and/or	[Model BOV-GP-STORA	GE-INTRA (7);]]	
	(2) □ and/or	[II.3.1.	semen coll production product procentre] (2) Commission regards real] (2) □ [P. Regulation	ection centre] (2) (6) n team] (2) (6) [and] (3 rocessing establishment] (6) situated in a third co on Implementing Regula sponsibilities, operation Part 2] (2) [Part 3] (2)	☐ [produced] (2), ☐ [processe [by an embryo collection tear 2) ☐ [processed] (2) ☐ [store (2) (6) ☐ [and stored in a gerountry or territory, or zone the tion (EU) 2021/404 and comple al procedures, facilities and e☐ [Part 4] (2) ☐ [Part 5] (2) or cred the Union accompanied by	n] (2) (6)
	(2)		\square either	[Model BOV-SEM-A-EN	TRY (7);]	
	(2)		□ and/or	[Model BOV-SEM-B-EN	TRY (7);]	
	(2)		\square and/or	[Model BOV-SEM-C-EN	ΓRY (7);]	
	(2)		□ and/or	[Model 1 in Part 1, Sect 2011/630/EU(7);]	ion A, of Annex II to Commiss	sion Implementing Decision
	(2)		\square and/or	[Model 2 in Part 1, Sect 2011/630/EU(7);]	ion B, of Annex II to Impleme	nting Decision
	(2)		\square and/or	[Model 3 in Part 1, Sect 2011/630/EU(7);]	ion C, of Annex II to Impleme	nting Decision
	(2)		\square and/or	[Model BOV-OOCYTES-	EMB-A-ENTRY(7);]	
	(2)		\square and/or	[Model BOV-in-vivo-EM	IB-B-ENTRY(7);]	
	(2)		\square and/or	[Model BOV-in-vitro-EN	MB-C-ENTRY(7);]	
	(2)		\square and/or	[Model BOV-in-vitro-EN	MB-D-ENTRY(7);]	
	(2)		\square and/or	[Model BOV-GP-PROCE	SSING-ENTRY(7);]	
	(2)		\square and/or	[Model BOV-GP-STORA	GE-ENTRY(7);]]	
		II.3.2.		=	l and stored in accordance wi to Delegated Regulation (EU) 2	
		II.3.3.	requireme	ents provided for in Artica), of Commission Delega	ckages on which the mark is a cle 10 of Delegated Regulation ated Regulation (EU) 2020/692	(EU) 2020/686 and/or Article
		II.3.4.	is/are trans	sported in a container w	hich:	
			II.3.4.1.	product storage centre	red prior to the date of dispat under responsibility of the co nd the seal bears the number	entre veterinarian, or by an
			II.3.4.2.	has been cleaned and e	either disinfected or sterilised	before use, or is single-use

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		2024/1044 (2021/403) MODEL BOV-GP-510RAGE-INTRA
II. Health info	ormation	
		container;
(2)(8)		☐ [II.3.4.3. has been filled in with a cryogenic agent which has not been previously used for other products;]
(2)(9)	□ [II.3.5.	is/are placed in straws or other packages which are securely and hermetically sealed;
ication	II.3.6.	is/are transported in a container where the different types are separated from each other by physical compartments or by being placed in secondary protective bags.]
Notes		
from the E Protocol or	uropean Ui n Ireland/N	te Agreement on the withdrawal of the United Kingdom of Great Britain and Northern Ireland nion and the European Atomic Energy Community, and in particular Article 5(4) of the orthern Ireland in conjunction with Annex 2 to that Protocol, references to the Union in this ate include the United Kingdom in respect of Northern Ireland.
provided f		rtificate shall be completed in accordance with the notes for the completion of certificates er 2 of Annex I to Commission Implementing Regulation (EU) 2020/2235.
Part I: Box reference I.11:	product st germinal referred t	dispatch": Indicate the unique approval number and the name and address of the germinal corage centre of dispatch of the consignment of semen, oocytes, and/or embryos. Only product storage centres approved by the competent authority and included in the register o in Article 101(1), point (b), of Regulation (EU) 2016/429 of the European Parliament and of il and Article 7 of Delegated Regulation (EU) 2020/686.
Box reference I.12:		destination": Indicate the address and unique registration or approval number of the nent of destination of the consignment of semen, oocytes, and/or embryos.
Box reference I.17:	to the seri accompar where the by which processing the germi product so	Inying documents": Number(s) of related original animal health certificate(s) shall correspond all number of the individual official document(s) or animal health certificate(s) that nied the semen, oocytes and/or embryos described in Part I from the semen collection centre semen was collected, and/or from the embryo collection team or the embryo production team the oocytes and/or embryos were collected or produced, and/or from the germinal product g establishment where the semen, oocytes or embryos were processed and stored, and/or from nal product storage centre where the semen, oocytes or embryos were stored, to the germinal corage centre described in box I.11. The original(s) of those document(s) or those animal health (s) or the officially endorsed copies thereof shall be attached to this animal health certificate.
Box reference I.19:	Seal numl	per shall be indicated.
Box reference I.26:	Total num	aber of packages shall correspond to the number of containers.
Box reference I.30:	"Species":	Select amongst "Bos taurus", "Bison bison" or "Bubalus bubalis" as appropriate.
		"Type": specify if semen, in vivo derived embryos, in vivo derived oocytes, in vitro produced embryos or micromanipulated embryos.
		"Identification number": Indicate identification number of each donor animal.
		ation mark": in Indicate mark on the straw or other packages where the semen, oocytes and/or of the consignment are placed.
		ollection/production": Indicate the date on which the semen, oocytes and/or embryos of the ent was/were collected or produced.
	of the sem	or registration number of plant/establishment/centre": Indicate the unique approval number nen collection centre where the semen of the consignment was collected, and/or of the embryo team or the embryo production team by which the oocytes or embryos of the consignment ected or produced.

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II. Health info Part II: (1) (2) (3) (4)	"Quantity": Indicate number of straws or other packages with the same mark. Only germinal product storage centres approved by the competent authority and included in the register referred to in Article 101(1), point (b), of Regulation (EU) 2016/429 and Article 7 of Delegated Regulation (EU) 2020/686. Delete if not applicable.							
(1) (2) (3)	Only germinal product storage centres approved by the competent authority and included in the register referred to in Article 101(1), point (b), of Regulation (EU) 2016/429 and Article 7 of Delegated Regulation (EU) 2020/686. Delete if not applicable.							
(1) (2) (3)	Only germinal product storage centres approved by the competent authority and included in the register referred to in Article 101(1), point (b), of Regulation (EU) 2016/429 and Article 7 of Delegated Regulation (EU) 2020/686. Delete if not applicable.							
(1) (2) (3)	register referred to in Article 101(1), point (b), of Regulation (EU) 2016/429 and Article 7 of Delegated Regulation (EU) 2020/686. Delete if not applicable.							
(2) (3)	register referred to in Article 101(1), point (b), of Regulation (EU) 2016/429 and Article 7 of Delegated Regulation (EU) 2020/686. Delete if not applicable.							
(3)								
(4)	Insert the name of the disease(s).							
ĺ	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)	Only germinal product establishments approved by the competent authority and included in the register referred to in Article 101(1), point (b), of Regulation (EU) 2016/429 and Article 7 of Delegated Regulation (EU) 2020/686.							
(7)	The original(s) of the document(s) or the animal health certificate(s) or the officially endorsed copies thereof that accompanied the semen, oocytes or embryos described in Part I from the semen collection centre where the semen was collected, and/or from the embryo collection team or the embryo production team by which the oocytes and/or embryos were collected or produced, and/or from the germinal product processing establishment where the semen, oocytes or embryos were processed and stored, and/or from the germinal product storage centre where the semen, oocytes or embryos were stored, to the germinal product storage centre of the semen, oocytes and/or embryos dispatch described in box I.11 shall be attached to this animal health certificate.							
(8)	Applicable for frozen semen, oocytes or embryos.							
(9)	Applicable for consignments where semen, oocytes, in vivo derived embryos, in vitro produced embryos and micromanipulated embryos of bovine animals are placed and transported in one container.							
Certifying Off Name (in cap Date of declar Stamp	·							

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