EUROPEAN UNION INTRA

ı	I.1. Consignor					I.2. IMSOC ref	erence	I.2.a. I.	ocal reference
	Name							-	ntral Competent Authority
	Address								cal Competent Authority
	Country ISO Code								,
Ļ	,							1	
:	I.5. Consignee					I.6. Operator conducting assembly operations independently of an establishment  Name			
	Name								
	Address					Address			
þ	Country			ISO Code		Address Approval Nu	mhor		
1						Country	itibei		ISO Code
3						Country	ountry 150 code		
;[	I.7. Country of orig	in			ISO Code	I.9. Country of	destination		ISO Code
il									
	I.O. Dogion of onigin					I 10 Dogion of	destination		Code
;†	I.1. Place of dispatch					I.10. Region of			Code
	_	tcn					iestination		
	Name					Name			
il	Address					Address			
:	Approval Number			*****		Approval Nu	mber		**************************************
	Country			ISO Code		Country			ISO Code
t	I.13. Place of loading	ng				I.14. Date and	time of departure		
l	Name	U					4		
l	Address								
l		,							
l	Approval Number					_ ^			
┨	Country ISO Code								
ſ	I.15. Means of Trai	nsport				I.16. Transpor	ter		
	Mode	Internation	al	Identification		Name			
		transport				Address			
ı		document				Approval Nu	mber		
l						Country			ISO Code
ŀ									
I						I.17. Accompa	nying documents		
						[en]			
				accompanyi ng.documen t.document.		Date o	ficena		
				t.document.					
					number		P.1	c	
						Country		Place o	Ĭ
ľ	I.18. Transport conditions					-1			
	Chilled						Ambient		
	crimeu 🗖			TTOZER	_	THISICIL L			
Ī	I.19. Container No / Seal No								
L									
	I.20. Certified as								
	Germinal products $\square$								
ŀ									
	Exit point					100 0 1	ISO Code		
						BCP code			
	I.22. For transit through Member State(s)				BCP code				
L					_	25. For export			
-							Third country ISO Code Exit point BCP code 1.25. Journey Log		
-	Member State								
	Member State	m ov tim o			I.24. Estimated journey time				
	Member State	rney time				1.20. journey	-06		
	Member State I.24. Estimated jou						-		
	Member State  I.24. Estimated jou  I.26. Total number	of packages				I.27. Total qua	-		
	Member State I.24. Estimated jou	of packages	:				-		
	Member State I.24. Estimated jou I.26. Total number I.28. Total gross we	of packages					-		
	Member State  I.24. Estimated jou  I.26. Total number  I.28. Total gross we  I.30. Description of	of packages eight f consignmen	nt			I.27. Total qua	ntity		
	Member State I.24. Estimated jou I.26. Total number I.28. Total gross we	of packages eight f consignmen		rs	Identification	I.27. Total qua	-		Nature of commodity
	Member State  I.24. Estimated jou  I.26. Total number  I.28. Total gross we  I.30. Description of	of packages eight f consignmen	nt	rs	Identification	I.27. Total qua	ntity		Nature of commodity
	Member State  I.24. Estimated jou  I.26. Total number  I.28. Total gross we  I.30. Description of	of packages eight f consignmen	nt	Package count	Identification	I.27. Total qua	ntity	Plant /	Nature of commodity  Establishment / Centre
	Member State I.24. Estimated jou I.26. Total number I.28. Total gross we I.30. Description of Commodity	of packages eight f consignmen	nt		Identification	I.27. Total qua	Quantity	Plant /	

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## **EUROPEAN UNION**

	** ** 1.1 · C				-						
	II. Health info	rmation									
Part II: Certification	I, the undersigned official veterinarian, hereby certify that:										
	II.1. The germinal product processing establishment(1) described in Box I.11. at which the semen(2)/										
		oocytes(2)/ in vivo derived embryos(2)/ in vitro produced embryos(2)/ micromanipluated embryos(2) was/were processed and stored:									
		II.1.1.	is approve	d and kept in a register l	by the competent authority;						
		II.1.2.			gards responsibilities, operational procedures, facilities Annex I to Commission Delegated Regulation (EU)						
	II.2.		en(2)/ oocytes(2)/ in vivo derived embryos(2)/ in vitro produced embryos(2)/ micromanipluated (2) described in Part I is/are intended for artificial reproduction and								
	(2) □ either	[II.2.1.	has/have been collected or produced, processed and stored in a semen collection centre(2)(3)/ by an embryo collection team(2)(3)/ by an embryo production team(2)(3), and/or processed and stored in a germinal product processing establishment(2)(3), and/or stored in a germinal product storage centre(2)(3) 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 processing establishment indicated in Box I.11. situated in the Member State of its/their collection or production under animal health certification requirements at least as strict as those provided for in:								
		(2)	□ either	[Model EQUI-SEM-A-IN							
		(2)	□ and/or	[Model EQUI-SEM-B-IN	TRA(4);]						
		(2)	$\square$ and/or	·							
		(2)	$\square$ and/or	[Model EQUI-SEM-D-IN	TRA(4);]						
		(2)	$\square$ and/or	[Model EQUI-OOCTYES	-EMB-A-INTRA(4);]						
		(2)	$\square$ and/or	[Model EQUI-OOCTYES	-EMB-B-INTRA(4);]						
		(2)	$\square$ and/or	[Model EQUI-OOCTYES	-EMB-C-INTRA(4);]						
		(2)	$\square$ and/or	[Model EQUI-OOCTYES	-EMB-D-INTRA(4);]						
		(2)	□ and/or	[Model EQUI-GP-PROCE	ESSING-INTRA(4);]						
		(2)	□ and/or	[Model EQUI-GP-STORA	AGE-INTRA(4);]]						
(2)   III.2.1. has/have been collected or produced, produced, and/or centre(2)(3)/ by an embryo collection team and/or processed and stored in a germinal stored in a germinal product storage centre collection or production and complying we operational procedures, facilities and equivalent of Annex I to Delegated Regular germinal product processing establishment Member State accompanied by certificate(		on team(2)(3)/ by an embryo erminal product processing ege centre(2)(3) situated in the lying with requirements as rond equipment set out in Part ed Regulation (EU) 2020/686, lishment indicated in Box I.1	production team(2)(3), stablishment(2)(3), and/or Member State of its/their egards responsibilities, 1(2)/Part 2(2)/Part 3(2)/Part and was/were moved to the 1. situated in another								
		(2)	$\square$ either	[Model EQUI-SEM-A-IN	TRA(4);]						
		(2)	$\square$ and/or	[Model EQUI-SEM-B-IN	TRA(4);]						
		(2)	$\square$ and/or	[Model EQUI-SEM-C-IN	TRA(4);]						
		(2)	$\square$ and/or	[Model EQUI-SEM-D-IN	TRA(4);]						
		(2)	$\square$ and/or	[Model EQUI-OOCTYES	-EMB-A-INTRA(4);]						
		(2)	$\square$ and/or	[Model EQUI-OOCTYES	-EMB-B-INTRA(4);]						
		(2)	$\square$ and/or	[Model EQUI-OOCTYES	-EMB-C-INTRA(4);]						
		(2)	$\square$ and/or	[Model EQUI-OOCTYES	-EMB-D-INTRA(4);]						
		(2)	$\square$ and/or	[Model EQUI-GP-PROCE							
		(2)	□ and/or	[Model EQUI-GP-STORA	AGE-INTRA(4);] ]						
	l										

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## **EUROPEAN UNION**

EC	IKOPEAN	ONION		ζΕŲ	JI-GP-PROCESSING-INTRA)
	II. Health inf	ormation			
Part II: Certification	(2) □ and/or	[II.2.1.	has/have been collected or produce centre(2)(3)/ by an embryo collectic and/or processed and stored in a gestored in a germinal product storage zone thereof listed in Annex XII to and complying with requirements facilities and equipment set out in to Delegated Regulation (EU) 2020/oin accordance with:	on team(2)(3)/ by an embryo perminal product processing e ge centre(2)(3) situated in a the Commission Implementing R as regards responsibilities, op Part 1(2)/Part 2(2)/Part 3(2)/Pa	production team(2)(3), stablishment(2)(3), and/or nird country, territory or egulation (EU) [C(2021)1800] perational procedures, art 4(2)/Part 5(2) of Annex I
ji,		(2)	□ either [Model EQUI-SEM-A-EN	TTRY(4);]	
  ii		(2)	☐ and/or [Model EQUI-SEM-B-EN		
art		(2)	☐ and/or [Model EQUI-SEM-C-EN	TRY(4);]	
P		(2)	☐ and/or [Model EQUI-SEM-D-EN		
		(2)	☐ and/or [Model EQUI-OOCYTES		
		(2)	☐ and/or [Model EQUI-OOCYTES	-EMB-B-ENTRY(4);]	
		(2)	☐ and/or [Model EQUI-OOCYTES	-EMB-C-ENTRY(4);]	
		(2)	☐ and/or [Model EQUI-GP-PROCE		
		(2)	☐ and/or [Model EQUI-GP-STORA		
		II.2.2.	has/have been collected, processed requirements set out in Annex III t		
		II.2.3.	is/are placed in straws or other pa requirements provided for in Artic 83(a) of Commission Delegated Reg I.30;	le 10 of Delegated Regulation	(EU) 2020/686 and/or Article
		II.2.4.	is/are transported in a container w	hich:	
			storage centre under re	ered prior to the dispatch from esponsibility of the centre vet eal bears the number as indic	erinarian, or by an official
			use container;	either disinfected or sterilised	d before use, or is a single-
		(2)(5)	has been filled in with for other products;]	the cryogenic agent which no	ot have been previously used
	(2)(6)	□ [II.2.5.	is/are placed in straws or other pa	ckages which are securely an	d hermetically sealed;
		II.2.6.	is/are transported in a container w compartments or by being placed i	, .	, ,

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## 2021/403 Equine germinal product from the processing establishment **EUROPEAN UNION** (EQUÍ-GP-PROCESSING-INTRA) II. Health information 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: Certification Box "Place of dispatch": Indicate the unique approval number and the name and address of the germinal reference product processing establishment of dispatch of the consignment of semen, oocytes, and/or embryos. I.11: Only germinal product processing establishments 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. Part II: "Place of destination": Indicate the address and unique registration or approval number of the Box reference establishment of destination of the consignment of semen, oocytes, and/or embryos. I.12: Box "Accompanying documents": Number(s) of related original certificate(s) shall correspond to the serial number of the individual official document(s) or health certificate(s) that accompanied the semen, reference oocytes and/or embryos described in Part I from the semen collection centre where the semen was collected, and/or the embryo collection and/or production team by which the oocytes and/or embryos were collected or produced, and/or the germinal product processing establishment where the semen, oocytes or embryos were processed and stored, and/or the germinal product storage centre where the semen, oocytes or embryos were stored to the germinal product processing establishment described in Box I.11. The original(s) of those document(s) or those certificate(s) or the officially endorsed copies thereof must be attached to this certificate. Seal number shall be indicated. Box reference I.19: Box Total number of packages shall correspond to the number of containers. reference I.26: "Type": Specify if semen, in vivo derived embryos, in vivo derived oocytes, in vitro produced embryos reference or micromanipulated embryos. I.30: "Identification number": Indicate identification number of each donor animal. "Identification mark": Indicate mark on the straw or other packages where semen, oocytes and/or embryos of the consignment are placed. "Date of collection/production": Indicate the date on which semen, oocytes and/or embryos of the consignment was/were collected or produced. "Approval or registration number of plant/establishment/centre": Indicate the unique approval number of the semen collection centre where the semen was collected, and/or of the embryo collection and/or production team by which the oocytes or embryos were collected or produced. "Quantity": Indicate number of straws or other packages with the same mark. Part II: (1) Only germinal product processing establishments approved by the competent authority and included in

- Only germinal product processing establishments 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.
- (2) Delete if not applicable.
- Only germinal product establishments 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.

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## **EUROPEAN UNION**

E	JROPEAN U	JNION	(EQUÍ-GP-PROCESSING-INTRA)					
	II. Health info	rmation						
Part II: Certification	(4)	that accompanied the semen, oocytes or embry where the semen was collected, and/or the em and/or embryos were collected or produced, at where the semen, oocytes or embryos were pr centre where the semen, oocytes or embryos w	h certificate(s) or the officially endorsed copies of thereof yos described in Part I from the semen collection centre bryo collection or production team by which the oocytes nd/or the germinal product processing establishment rocessed and stored, and/or the germinal product storage were stored to the germinal product processing abryos dispatch described in Box I.11 must be attached to					
	(5)	Applicable for frozen semen, oocytes or embryos.						
	(6)	Applicable for the consignment where in one container semen, oocytes, in vivo derived embryos, in						
	vitro produced embryos and micromanipulated embryos of equine animals are placed and transported.							
	Certifying Offi	icer/Official veterinarian	0.100					
	Name (in cap Date of signa		Qualification and title Signature					
	Stamp							

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