EUROPEAN UNION

	I.1. Consignor					I.2. IMSOC ref	erence	I.2.a. Local reference		
	Name						I.3. Central Competen	I.3. Central Competent Authority		
	Address							I.4. Local Competent	Authority	
	Country		ISO Co	de						
님	I.5. Consignee					I.6. Operator conducting assembly operations independently of an establishment				
Part I: Description of consignment	Name						L .			
놀	Address					Name				
<u>છ</u>	Country		ISO Co	de		Address Approval Number Country ISO Code				
ns										
ខ										
Ч	I.7. Country of orig	gin			ISO Code	I.9. Country of	f destination		ISO Code	
g	, , ,					, , , , , , , , , , , , , , , , , , ,				
B										
믭	I.8. Region of origi				Code	I.10. Region of			Code	
ទ្ធ	I.11. Place of dispa	itch				I.12. Place of destination				
ë	Name					Name				
-	Address					Address Approval Number Country ISO Code				
ย่	Approval Number	r								
Pa	Country		ISO	Code						
ł	I.13. Place of loadi	ng				I.14. Date and time of departure				
		••o				1.14. Date and time of departure				
	Name									
	Address									
	Approval Number	r		. ·						
	Country		ISO	Code						
	I.15. Means of Trai	nsport				I.16. Transpor	ter			
	Mode	International	Identificati	on		Name				
	woue	transport	identificati	.011		Address				
		document				Activity ID				
						Country ISO Code				
						I.17. Accompanying documents				
						Commercial				
						document Date of issue reference				
						Countwr		Place of		
						Country issue				
	I.18. Transport cor	nditions		_	_		_			
	Ambient 🗆			Chilled [Frozen 🗆				
ŀ	I.19. Container No	/ Cool No								
	1.19. Container No	/ Sear NO								
Ī	I.20. Certified as									
	Germinal products									
	I.21. For transit th	rough a third c	ountry							
	Third country					ISO Code				
	Exit point					BCP code				
	Entry point					BCP code				
	I.22. For transit th	rough Member				I.23. For export				
	Member State ISO Code .26. Total number of packages I.27. Total quantity				Third countryISO CodeExit pointBCP code					
					I.25. Journey l					
					I.28. Total gross weight					
ŀ	L30. Description of	f consignment		1						
	I.30. Description of consignment Commodity Species Identification I					Number	Quantity	Notire of a	modity	
						munner	Quantity	Nature of com	mouly	
						. ,				
	Identification Ma	dentification Mark Package count Date of coll production		Date of collect	tion / Plant / Establishment / Centre		/ Туре			
					production					
ŀ	<u>L</u>					<u> </u>	I			
1										

EUROPEAN UNION

ΞU									
	II. Health inf	ormation							
		I, the unde	rsigned off	icial veterinarian, hereb	y certify that:				
	II.1.			s(1)/ embryos(1) describe he donor animals which	d in Part I are intended for ar	tificial reproduction and			
		II.1.1.							
Certification		II.1.2.	have remained in a single establishment of origin for a period of at least 30 days prior to the date of collection of the semen(1)/ oocytes(1)/ embryos(1);						
Cerun	(1)	□ [II.1.3.		als of the family Camelid on Delegated Regulation	ae and are identified in accordance with Article 73(1) of				
	(1)	□ [II.1.3.	are animals of the family Cervidae and are identified in accordance with Article 7 Article 74 of Delegated Regulation (EU) 2019/2035.]						
2	II.2.		(1)/ oocytes(1)/ embryos(1) described in Part I come from a registered establishment assigne petent authority with a unique registration number as indicated in Box I.11.						
	II.3.	According animals w		nformation, the semen(1)/ oocytes(1)/ embryos(1) were	e obtained from the donor			
	-	II.3.1.	establishr mouth dis infection	nent, situated in a restric sease, infection with rind	it, nor have been in contact w ted zone established due to th erpest virus, infection with Ri inants virus or of an emergin imals;	e occurrence of foot-and- ft Valley fever virus,			
		II.3.2.			e during a period of at least th e semen(1)/ oocytes(1)/ embry				
			II.3.2.1.	complex (M. bovis, M. o	nme to detect infection with M caprae and M. tuberculosis) h 2 or 3 of Annex II to Commissi	as been carried out in			
			II.3.2.2.		ily Camelidae or Cervidae whi to in point II.3.2.1. has been i				
			II.3.2.3.		infection with Mycobacteriur I. tuberculosis), investigations				
		II.3.3.	Brucella s	uis has not been reporte	e infection with Brucella abor d during the period of at least en(1)/ oocytes(1)/ embryos(1);				
	(1)	□ [II.3.4.	present ha and Bruce with nega	ave been subjected to a te ella suis as referred to in tive results carried out o	ae and come from an establish est for infection with Brucella Part 1 of Annex I to Delegated n samples taken during the po of the semen(1)/ oocytes(1)/ e	abortus, Brucella melitensis Regulation (EU) 2020/688 eriod of the preceding 30			
		II.3.5.	vulvovagi	nitis has not been report	e infectious bovine rhinotracl ed during the period of at leas e semen(1)/ oocytes(1)/ embry	st the preceding 30 days			
		II.3.6.	not been i	reported during a period of the semen(1)/ oocytes	e infection with epizootic hae of at least the preceding 2 yea (1)/ embryos(1) within a radiu	ars prior to the date of			
		II.3.7.	during the		e infection with rabies virus h eceding 30 days prior to the d				
		II.3.8.		preceding 15 days prior to	e anthrax has not been report o the date of collection of the s				

EUROPEAN UNION

	TT TT 1/1. 1. C.								
	II. Health info	ormation							
		II.3.9.	during a p		e surra (Trypanosoma evansi) has not been reported reding 30 days prior to the date of collection of the and				
	(1)	\circ either	[surra has	not been confirmed dur	ing the preceding 2 years;]				
lon	(1)	∘ or			the preceding 2 years and following the last outbreak of remained under movement restrictions until rere removed from the establishment; and				
cau			-	the infected animals w					
Part II: Ceruncation			-	(Trypanosoma evansi) (EU) 2020/688, with neg	on the establishment were su referred to in Part 3 of Annex gative result, carried out on sa ed animals were removed fro	I to Delegated Regulation Imples taken at least 6			
ra.		II.3.10.		th at least one of the foll otypes 1-24):	owing conditions as regards in	nfection with bluetongue			
	(1)	□ either	[II.3.10.1.	of the semen(1)/ oocyte from infection with blu	r a period of at least 60 days p es(1)/ embryos(1) in a Member letongue virus (serotypes 1-24 (serotypes 1-24) has been cont animal population;]	State or zone thereof free where no case of infection			
	(1)	□ and/or	[II.3.10.2.						
	(1)	□ and/or	[II.3.10.3.	disease-free period, for of the semen(1)/ oocyte where the competent a semen(1)/ oocytes(1)/ e competent authority of	a seasonally disease-free zon a period of at least 60 days pres(1)/ embryos(1), in a Member authority of the place of origin mbryos(1) has obtained the present the Member State of destinat easonally disease-free zone ar / embryos(1);]	rior to and during collection r State or zone thereof of the consignment of rior written consent of the ion to the conditions for			
	(1)	□ and/or	[II.3.10.4.		n a vector-protected establishr uring collection of the semen(1				
	(1)	□ and/or	[II.3.10.5.	bluetongue virus serog	ted to a serological test to dete roup 1-24, with negative resu ollection of the semen(1)/ ooc	lts, between 28 and 60 days			
	(1)	□ and/or	[II.3.10.6.	(serotypes 1-24), with r commencement and fir semen at intervals of a	ed to an agent identification t negative results, on blood sam nal collection of the semen an t least every 7 days, in the cas lays, in the case of PCR;]	ples taken at d during collection of the			
	(1)	□ and/or	[II.3.10.7.		ted to an agent identification negative results, on blood sam s(1)/ embryos(1).]				
	II.4.			wledge and as declared b re obtained from the doi	by the operator, the semen(1)/ nor animals which	oocytes(1)/ embryos(1)			
		II.4.1.		clinically examined by a ection of the semen(1)/ c	a veterinarian and showed no ocytes(1)/ embryos(1);	diseases symptoms on the			
		II.4.2.	in point II.		nals which did not comply wit to II.3.10. during the residency				
		II.4.3.		-	g during a period of at least 30 (1)/ embryos(1) and during the				

2021/403 Germinal products of camelidae and cervidae (GP-CAMELID-CERVID-INTRA)

	II. Health info	rmation							
	II.5.	The semen(1)/ oocytes(1)/ embryos(1) described in Part I are placed in a sealed transport container and the seal bears the number as indicated in Box I.19.							
	II.6.								
iffici	Notes								
Part II: (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 address and the unique registration number of the establishmer erence dispatch of the consignment of semen, oocytes or embryos.							
	Box reference I.12:	"Place of destination": Indicate the address and the unique registration number of the establishment of destination of the consignment of semen, oocytes or embryos.							
	Box reference I.30:	"Type": Specify if semen, in vivo derived embryos, in vivo derived oocytes, in vitro produced embryos or micromanipulated embryos.							
		"Species": Indicate "Camelidae" or "Cervidae"	as appropriate.						
		"Identification number": Indicate individual identification number of each donor animal.							
		"Identification mark": Indicate mark on the straw or other packages where semen, oocytes of the consignment are placed.							
		"Date of collection/production: Indicate the date on which semen, oocytes or embryos of the consignment were collected or produced.							
		"Approval or registration number of plant/establishment/centre": Indicate the unique registration number of the establishment of the collection or production of semen, oocytes or embryos of the consignment.							
	"Quantity": Indicate number of straws or other packages with the same mark.								
	Part II:								
	(1)	Delete if not applicable.							
	Name (in cap	cer/Official veterinarian	Authority name						
	Date of signa Stamp		Signature						

EUROPEAN UNION