

**ANIMAL HEALTH CERTIFICATE FOR THE MOVEMENT BETWEEN MEMBER STATES OF
CONSIGNMENTS OF SEMEN, OOCYTES AND EMBRYOS OF ANIMALS OF THE FAMILIES CAMELIDAE AND
CERVIDAE WHICH WERE COLLECTED OR PRODUCED, PROCESSED AND STORED IN ACCORDANCE WITH
REGULATION (EU) 2016/429 AND DELEGATED REGULATION (EU) 2020/686**

(MODEL 'GP-CAM-CER-INTRA')

EUROPEAN UNION		INTRA	
Part I: Description of consignment	I.1 Consignor	I.2 IMSOC reference	QR CODE
	Name	I.2a Local reference	
	Address	I.3 Central Competent Authority	
	Country ISO country code	I.4 Local Competent Authority	
	I.5 Consignee	I.6 Operator conducting assembly operations independently of an establishment	
	Name	Name	Registration No
	Address	Address	
	Country ISO country code	Country	ISO country code
	I.7 Country of origin ISO country code	I.9 Country of destination	ISO country code
	I.8 Region of origin Code	I.10 Region of destination	Code
	I.11 Place of dispatch	I.12 Place of destination	
	Name Registration/Approval No	Name	Registration/Approval No
	Address	Address	
Country ISO country code	Country	ISO country code	
I.13 Place of loading	I.14 Date and time of departure		
I.15 Means of transport	I.16 Transporter		
<input type="checkbox"/> Vessel <input type="checkbox"/> Aircraft	Name	Registration/Authorisation No	
<input type="checkbox"/> Railway <input type="checkbox"/> Road vehicle	Address		
Identification <input type="checkbox"/> Other	Country	ISO country code	
Document	I.17 Accompanying documents		
	Type	Code	
	Country	ISO country code	
	Commercial document reference		
I.18 Transport conditions <input type="checkbox"/> Ambient <input type="checkbox"/> Chilled <input type="checkbox"/> Frozen			
I.19 Container number/Seal number			
Container No	Seal No		

I.20 Certified as or for							
<input type="checkbox"/> Further keeping	<input type="checkbox"/> Slaughter	<input type="checkbox"/> Confined establishment	<input type="checkbox"/> Germinal products				
<input type="checkbox"/> Registered equine animal	<input type="checkbox"/> Travelling circus/animal act	<input type="checkbox"/> Exhibition	<input type="checkbox"/> Event or activity near borders				
<input type="checkbox"/> Release into the wild	<input type="checkbox"/> Dispatch centre	<input type="checkbox"/> Relaying area/purification centre	<input type="checkbox"/> Ornamental aquaculture establishment				
<input type="checkbox"/> Further processing	<input type="checkbox"/> Organic fertilizers and soil improvers	<input type="checkbox"/> Technical use	<input type="checkbox"/> Quarantine or similar establishment				
<input type="checkbox"/> Products for human consumption	<input type="checkbox"/> Pollination	<input type="checkbox"/> Live aquatic animals for human consumption	<input type="checkbox"/> Other				
I.21 <input type="checkbox"/> For transit through a third country							
Third country		ISO country code					
Exit point		BCP code					
Entry point		BCP code					
I.22 <input type="checkbox"/> For transit through Member State(s)				I.23 <input type="checkbox"/> For export			
Member State	ISO country code	Third country	ISO country code				
Member State	ISO country code	Exit point	BCP code				
Member State	ISO country code						
I.24 Estimated journey time				I.25 Journey log <input type="checkbox"/> yes <input type="checkbox"/> no			
I.26 Total number of packages				I.27 Total quantity			
I.28 Total net weight/gross weight (kg)				I.29 Total space foreseen for the consignment			
I.30 Description of consignment							
CN code	Species	Subspecies/Category	Sex	Identification system	Identification number	Age	Quantity
							Type
Region of origin		Cold store		Identification mark	Type of packaging		Net weight
Slaughterhouse		Treatment type		Nature of commodity	Number of packages		Batch No
		Date of collection/production		Manufacturing plant	Approval or registration number of plant/establishment/centre	Test	

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Certificate model GP-CAMELID-CER-INTRA

Part II: Certification	II. Health information	II.a Certificate reference	II.b IMSOC reference
		<p>I, the undersigned official veterinarian, hereby certify that:</p> <p>II.1. The semen⁽¹⁾/ oocytes⁽¹⁾/ embryos⁽¹⁾ described in Part I are intended for artificial reproduction and were obtained from donor animals which</p> <p>II.1.1. have been born and remained since birth in the Union, or have entered the Union in accordance with the requirements for entry into the Union;</p> <p>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⁾;</p> <p>⁽¹⁾II.1.3. are animals of the family Camelidae and are identified in accordance with Article 73(1) of Commission Delegated Regulation (EU) 2019/2035.]</p> <p>⁽¹⁾II.1.3. are animals of the family Cervidae and are identified in accordance with Article 73 (2) or Article 74 of Delegated Regulation (EU) 2019/2035.]</p> <p>II.2. The semen⁽¹⁾/ oocytes⁽¹⁾/ embryos⁽¹⁾ described in Part I comes/come from a registered establishment assigned by the competent authority with a unique registration number as indicated in Box I.11.</p> <p>II.3. According to official information, the semen⁽¹⁾/ oocytes⁽¹⁾/ embryos⁽¹⁾ was/were obtained from donor animals which</p> <p>II.3.1. do not come from an establishment, nor have been in contact with animals from an establishment, situated in a restricted zone established due to the occurrence of foot-and-mouth disease, infection with rinderpest virus, infection with Rift Valley fever virus, infection with peste des petits ruminants virus or of an emerging disease relevant for species of those kept terrestrial animals;</p> <p>II.3.2. come from an establishment where during a period of at least the preceding 12 months prior to the date of collection of the semen⁽¹⁾/ oocytes⁽¹⁾/ embryos⁽¹⁾</p> <p>II.3.2.1. a surveillance programme to detect infection with <i>Mycobacterium tuberculosis</i> complex (<i>M. bovis</i>, <i>M. caprae</i> and <i>M. tuberculosis</i>) has been carried out in accordance with Part 2 or 3 of Annex II to Commission Delegated Regulation (EU) 2020/688;</p> <p>II.3.2.2. no animals of the family Camelidae or Cervidae which do not fulfil the requirements referred to in point II.3.2.1. has been introduced;</p> <p>II.3.2.3. in case of suspicion of infection with <i>Mycobacterium tuberculosis</i> complex (<i>M. bovis</i>, <i>M. caprae</i> and <i>M. tuberculosis</i>), investigations were carried out and the disease was ruled out;</p> <p>II.3.3. come from an establishment where infection with <i>Brucella abortus</i>, <i>Brucella melitensis</i> and <i>Brucella suis</i> has not been reported during the period of at least the preceding 42 days prior to the date of collection of the semen⁽¹⁾/ oocytes⁽¹⁾/ embryos⁽¹⁾;</p>	

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Certificate model GP-CAMELID-CER-INTRA

	<p>⁽¹⁾[II.3.4. are animals of the family Camelidae and come from an establishment where all animals present have been subjected to a test for infection with <i>Brucella abortus</i>, <i>Brucella melitensis</i> and <i>Brucella suis</i> as referred to in Part 1 of Annex I to Delegated Regulation (EU) 2020/688 with negative results carried out on samples taken during the period of the preceding 30 days prior to the date of collection of the semen⁽¹⁾/ oocytes⁽¹⁾/ embryos⁽¹⁾];</p> <p>II.3.5. come from an establishment where infectious bovine rhinotracheitis/ infectious pustular vulvovaginitis has not been reported during the period of at least the preceding 30 days prior to the date of collection of the semen⁽¹⁾/ oocytes⁽¹⁾/ embryos⁽¹⁾;</p> <p>II.3.6. come from an establishment where infection with epizootic haemorrhagic disease virus has not been reported during a period of at least the preceding 2 years prior to the date of collection of the semen⁽¹⁾/ oocytes⁽¹⁾/ embryos⁽¹⁾ within a radius of 150 km around the establishment;</p> <p>II.3.7. come from an establishment where infection with rabies virus has not been confirmed during the period of at least the preceding 30 days prior to the date of collection of the semen⁽¹⁾/ oocytes⁽¹⁾/ embryos⁽¹⁾;</p> <p>II.3.8. come from an establishment where anthrax has not been reported during the period of at least the preceding 15 days prior to the date of collection of the semen⁽¹⁾/ oocytes⁽¹⁾/ embryos⁽¹⁾;</p> <p>II.3.9. come from an establishment where surra (<i>Trypanosoma evansi</i>) has not been reported during a period of at least the preceding 30 days prior to the date of collection of the semen⁽¹⁾/ oocytes⁽¹⁾/ embryos⁽¹⁾, and</p> <p>⁽¹⁾either [surra has not been confirmed during the preceding 2 years;]</p> <p>⁽¹⁾or [surra has been confirmed during the preceding 2 years and following the last outbreak of that disease the establishment has remained under movement restrictions until</p> <ul style="list-style-type: none"> – the infected animals were removed from the establishment; and – the remaining animals on the establishment were subjected to a test for surra (<i>Trypanosoma evansi</i>) referred to in Part 3 of Annex I to Delegated Regulation (EU) 2020/688, with negative result, carried out on samples taken at least 6 months after the infected animals were removed from the establishment;] <p>II.3.10. comply with at least one of the following conditions as regards infection with bluetongue virus (serotypes 1-24):</p> <p>⁽¹⁾either [II.3.10.1. they have been kept for a period of at least 60 days prior to and during collection of the semen⁽¹⁾/ oocytes⁽¹⁾/ embryos⁽¹⁾ in a Member State or zone thereof free from infection with bluetongue virus (serotypes 1-24) where no case of infection with bluetongue virus (serotypes 1-24) has been confirmed during the last 24 months in the targeted animal population;]</p> <p>⁽¹⁾and/or [II.3.10.2. they have been kept in a seasonally disease-free zone, during the seasonally disease-free period, for a period of at least 60 days prior to and during collection of the semen⁽¹⁾/ oocytes⁽¹⁾/ embryos⁽¹⁾, in a Member State or zone thereof with an approved eradication programme against infection with bluetongue virus (serotypes 1-24);]</p>
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EUROPEAN UNION

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	<p>⁽¹⁾and/or [II.3.10.3. they have been kept in a seasonally disease-free zone, during the seasonally disease-free period, for a period of at least 60 days prior to and during collection of the semen⁽¹⁾/ oocytes⁽¹⁾/ embryos⁽¹⁾, in a Member State or zone thereof where the competent authority of the place of origin of the consignment of semen⁽¹⁾/ oocytes⁽¹⁾/ embryos⁽¹⁾ has obtained the prior written consent of the competent authority of the Member State of destination to the conditions for establishment of that seasonally disease-free zone and to accept the consignment of semen⁽¹⁾/ oocytes⁽¹⁾/ embryos⁽¹⁾];</p> <p>⁽¹⁾and/or [II.3.10.4. they have been kept in a vector-protected establishment for a period of at least 60 days prior to and during collection of the semen⁽¹⁾/ oocytes⁽¹⁾/ embryos⁽¹⁾];</p> <p>⁽¹⁾and/or [II.3.10.5. they have been subjected to a serological test to detect antibodies to the bluetongue virus serogroup 1-24, with negative results, between 28 and 60 days from the date of each collection of the semen⁽¹⁾/ oocytes⁽¹⁾/ embryos⁽¹⁾];</p> <p>⁽¹⁾and/or [II.3.10.6. they have been subjected to an agent identification test for bluetongue virus (serotypes 1-24), with negative results, on blood samples taken at commencement and final collection of the semen and during collection of the semen at intervals of at least every 7 days, in the case of the virus isolation test, or of at least every 28 days, in the case of PCR;]</p> <p>⁽¹⁾and/or [II.3.10.7. they have been subjected to an agent identification test for bluetongue virus (serotypes 1-24), with negative results, on blood sample taken on the day of collection of the oocytes⁽¹⁾/ embryos⁽¹⁾.]</p> <p>II.4. To the best of my knowledge and as declared by the operator, the semen⁽¹⁾/ oocytes⁽¹⁾/ embryos⁽¹⁾ described in Part I was/were obtained from donor animals which</p> <p>II.4.1. have been clinically examined by a veterinarian and showed no disease symptoms on the day of collection of the semen⁽¹⁾/ oocytes⁽¹⁾/ embryos⁽¹⁾;</p> <p>II.4.2. have not been in contact with animals which did not comply with the requirements set out in point II.1.1. and in points II.3.1. to II.3.10. during the residence period of at least 30 days set out in point II.1.2.;</p> <p>II.4.3. were not used for natural breeding during a period of at least 30 days prior to the date of collection of the semen⁽¹⁾/ oocytes⁽¹⁾/ embryos⁽¹⁾ and during the collection period.</p> <p>II.5. The semen⁽¹⁾/ oocytes⁽¹⁾/ embryos⁽¹⁾ described in Part I is/are placed in a sealed transport container and the seal bears the number as indicated in Box I.19.</p> <p>II.6. To the best of my knowledge and based on the documentary check of the data submitted by the operator, the semen⁽¹⁾/ oocytes⁽¹⁾/ embryos⁽¹⁾ described in Part I is/are placed in straws or other packages on which the mark is applied in accordance with requirements provided for in Article 11 of Commission Delegated Regulation (EU) 2020/686 and that mark is indicated in Box I.30.</p>
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