						I.2. IMSOC Reference		
	Name				I.2.a. Local Reference			
	Address							
	Country		ISO Code					
	I.5. Consignee				I.3. Central co	mpetent authority		
	Name					petent authority		
en	Address							
gnm	Country		ISO Code					
Part I : Details of consignment	I.7. Country of orig	gin		ISO Code	I.9. Country of	f destination		ISO Code
fc	I.8. Region of origi	n		Code	I.10. Region of	destination		
S 0	I.11. Place of Dispa			coue	I.12. Place of c			
ail	Name				Name			
Det	Address				Address			
	Approval Number	r			Approval Nu	mber		
τI	Country		ISO Code		Country		ISO Code	
Par	-							
-	I.13. Place of Load	ing			I.14. Date and	time of departure		
	Name							
	Address							
	Approval Number	ſ	ISO Code					
	Country		ISO Code					
	I.15. Means of Trai	nsport			I.16 Entry Point			
	Mode	International	Identification					
		transport document						
	I.18. Transport cor	ditions			L17. Accompa	nying documents		
	Ambient 🗆				Commercial			
					document Date of issue			
					Diago of			
					Country	issu		
	I.19. Container No / Seal No							
	I.20. Certified as							
	Registered equidae	e 🗆						
	I.21. For transit th	rough a third cou	ntry 🗌		I.22. For transit through Member State(s)			
	Country		ISO Code		Country ISO Code			
	EU Exit Authority		BCP code					
	EU Entry		BCP code					
	Authority I.25. Total gross we	eight	201 0040					
	1.25. 10tal gross we	eigne						
	-	-						
	I.28. Description of	f consignment						
	I.28. Description of 1.01 LIVE ANIMA	f consignment	nd hinnies					
	I.28. Description of 1.01 LIVE ANIMA 0101 Live horse	f consignment LS es, asses, mules ar		Breed/Categor	۳V	Identification mark	Identification	number
	I.28. Description of 1.01 LIVE ANIMA	f consignment LS		Breed/Categor	су	Identification mark	Identification	number
	I.28. Description of 1. 01 LIVE ANIMA 0101 Live horse Commodity	f consignment LS es, asses, mules ar		Breed/Categor	1	Identification mark	Identification	number
	I.28. Description of 1.01 LIVE ANIMA 0101 Live horse	f consignment LS es, asses, mules ar		Breed/Categor	ry Gender	Identification mark	Identification	number
	I.28. Description of 1. 01 LIVE ANIMA 0101 Live horse Commodity	f consignment LS es, asses, mules ar		Breed/Categor	1	Identification mark	Identification	number
	I.28. Description of 1. 01 LIVE ANIMA 0101 Live horse Commodity	f consignment LS es, asses, mules ar		Breed/Categor	1	Identification mark	Identification	number
	I.28. Description of 1. 01 LIVE ANIMA 0101 Live horse Commodity	f consignment LS es, asses, mules ar		Breed/Categor	1	Identification mark	Identification	number
	I.28. Description of 1. 01 LIVE ANIMA 0101 Live horse Commodity	f consignment LS es, asses, mules ar		Breed/Categor	1	Identification mark	Identification	number
	I.28. Description of 1. 01 LIVE ANIMA 0101 Live horse Commodity	f consignment LS es, asses, mules ar		Breed/Categor	1	Identification mark	Identification	number
	I.28. Description of 1. 01 LIVE ANIMA 0101 Live horse Commodity	f consignment LS es, asses, mules ar		Breed/Categor	1	Identification mark	Identification	number
	I.28. Description of 1. 01 LIVE ANIMA 0101 Live horse Commodity	f consignment LS es, asses, mules ar		Breed/Categor	1	Identification mark	Identification	number
	I.28. Description of 1. 01 LIVE ANIMA 0101 Live horse Commodity	f consignment LS es, asses, mules ar		Breed/Categor	1	Identification mark	Identification	number

II. Health info	rmation		_					
			ا ficial veterinarian, hereby requirements:	certify that the equine	animal(s)	described above		
	II.1	it/they co	ome(s) from a Member Stat	e of the European Unio	n:			
		II.1.1	in which African horse encephalomyelitis, equi and dourine (Trypanose	ine infectious anaemia,	glanders	(Burkholderia mallei)		
		II.1.2	that is considered by th encephalitis and Venez restrictive measures ar State described in Box I compliance with all rele	uelan equine encephalo e in place on these disea .7., and the Member Sta	myelitis a ises by th te descrik	and in which no e EU or the Member bed in Box I.7. is in full		
		II.1.3		hada and in which no re J or the Member State d d in Box I.7. is in full cor	strictive escribed			
	II.2	during the 6 months immediately prior to export to Canada, it/they has/have not been in any country or zone in which Venezuelan equine encephalomyelitis has occurred in the past 24 months, it/they has/have not been vaccinated against Venezuelan equine encephalomyelitis within 60 days of export to Canada, and the Member State described in Box I.7. is in full compliance with all relevant EU legislation for this disease;						
	II.3	it/they has/have been continually resident in the EU for a minimum of 60 days, or since birth if less than 60 days of age, immediately preceding the pre-export isolation certified in point II.7 for export to Canada;						
	II.4	during the 90 days immediately prior to export to Canada, it/they has/have not been in contact with equidae (including imported horses) that have been in an area where restrictive measures are in place on African horse sickness or in a country or zone where African horse sickness has been diagnosed in the past 60 days, and it/they has/have not been vaccinated against African horse sickness within 60 days of export to Canada, and the Member State described in Box I.7. is in full compliance with all relevant EU legislation for this disease;						
	II.5	during the 90 days immediately prior to export to Canada, it/they has/have not been on any premises subject to restrictive measures for glanders or dourine and it/they has/have not had contact with equidae (including imported horses) that have been in an area where restrictive measures are in place on dourine and glanders during the past 6 months and the Member State described in Box I.7., is in full compliance with all relevant EU legislation for these diseases;						
	II.6	during the 90 days immediately prior to export to Canada, it/they has/have not been premises where equine piroplasmosis (Theileria equi and Babesia caballi) or equine infectious anaemia has occurred nor has equine infectious anaemia occurred on any adjoining premises;						
	II.7	requiren veterina describe	ns/have been isolated for th nents, immediately prior to rian officially recognised b d in Box I.7, and it/they has us disease during that isola	export to Canada on a y the competent author /have remained free fro	premises ity of the	approved by a EU Member State		
	II.8		ne isolation period immedi as/have had blood samples					
		II.8.1	for equine infectious ar alternate test acceptabl	-	or, wher	e applicable, an		
		II.8.2	for equine piroplasmos where applicable, an al has/have been maintair treatment, during the 3	ternate test acceptable t ned free from ticks, whe	o CFIA; a n necessa	nd the animal(s)		

Part II: Certification

II. Health information								
II.9	II.9 during the 90 days immediately preceding exportation to Canada the animal(s) I been on a premises where contagious equine metritis (CEM) has occurred							
	no manipulation or treatment of the reproductive tract, except collection of swabs where required, has been performed during the 30 days preceding exportation;							
and	(1)either		mal(s) is/are ation comme		ays of age(2) on the day pre-			
		and	have never been bred nor has breeding of the horse(s) been attempted and it/they has/have never been commingled and left unattended with adult equidae of the opposite sex, except in case of foals left with their dam					
		and	relevant po	er or his/her representative has/have been advised of the post-import conditions that must be met, as outlined in the n Import Permit(3),				
		and		quirements in points II.10 and II.11 for contagious equine EM) do not apply.]				
	(1)or		mal(s) is/are isolation co		er 731 days of age on the day			
		and	relevant po	the owner or his/her representative has/have been advised of the relevant post-import conditions that must be met, as outlined in the Canadian Import Permit(3),				
		and	described in samples tak specimens I the supervi	ten within the 30 days pri have been collected(4)(5) sion of an official vetering 48 hours of collection in	ce with the procedure and in point II.11 for mares with or to export, in which case all by a licensed veterinarian under arian and were cultured for a laboratory officially approved			
II.10	stallion(s), in the country of origin, during the 30 days preceding exportation has/have n been mated by natural breeding or has/have not had semen collected for the purpose of artificial insemination and within that same period one (1) set of three (3) specimens (swabs) has been collected from the prepuce (sheath), the fossa glandis (same as urethra fossa) including the diverticulum (same as the urethral sinus) and the terminal (distal) e of the urethra, and all specimens were subjected to the required test for CEM(6)(7) with							
(1)	either	○ [negativ	ve results as s	specified in the table in po	int II.12 below;]			
(1)	or	• [negative results obtained on specimens taken not less than 21 days after the completion of the treatment of the stallion(s) for CEM carried out in a manner approved by the competent authority of the EU Member State following a positive result in a previous test for CEM as specified in the table in point II.12 below and the stallion(s) has/have been test mated to two mares in each case which have been subjected with negative results to						
		-	set of three after matin lateral and	(3) specimens (swabs), co g, from the mucosal surfa medial clitorial sinuses, a ım instead of the cervix, i	y culture carried out on one (1) llected not earlier than 3 days ces of the clitorial fossa, the nd the cervix (or the n the case the mare(s) is/are in			
		-		ent fixation test for the de equigenitalis carried out o	tection of antibodies to n samples taken 21-30 days post			
II.11 mare(s), in the country of origin, during the 30 days preceding exportation been mated by natural breeding nor artificial insemination and within t								

Part II: Certification

			•		L		
	II. Health information						
	(1)	either	 • [the mare(s) is/are not pregnant and one (1) set of three (3) specimens (swabs) has been collected from the mucosal surfaces of the clitorial fossa, the lateral and medial clitorial sinuses, and the cervix, (or the endometrium instead of the cervix in the case the mare(s) is/are in oestrus);] 				
TOTIOTI	(1)	or	 [the mare(s) is/are pregnant and one (1) set of two (2) specimens (swabs) has been collected from the mucosal surfaces of the clitorial fossa and the lateral and medial clitorial sinuses (swabbing of the cervix and endometrium do not apply);] 				
		and	all specime	ens were sul	pjected to the required test f	or CEM(6) (7) with:	
Ś	(1)	either	○ [negativ	egative results as specified in the table in point II.12 below]			
	(1)	or	on specime of the mar authority of for CEM as been subje	ens taken no e(s) for CEM of the EU Me specified in cted with no	ot less than 21 days after the carried out in a manner ap mber State following a posi	ive result in a previous test w, and the mare(s) has/have ment fixation test for the	
		and	(1)either	artificially pre-export	re(s) has/have not been mate inseminated within the last period during which no bre on is allowed;]	21 days preceding the 30-day	
			(1)or	preceding artificial in negative re	the 30-day pre-export period semination is allowed and l sult to a complement fixation aken between 21 and 30 day	ly inseminated within 21 days l during which no breeding or nas/have been subjected with on test, carried out on (a) blood ys after artificial	
II.12 Details (7) or			on testing and treatments for CEM as referred to in points II.10 and II.11				
	Date and time of specimen collection (A)	0	Results (C)	Name of the official laboratory (D)			

II. Health inf	Formation II.13	it/they has/have been inspected on (dd/mm/yyyy) within 72 hours prior to			
	II.13	it/they has/have been inspected on (dd/mm/yyyy) within 72 hours prior to			
		loading for export to Canada by a veterinarian officially recognised by the competent authority of the EU Member State described in Box I.7 and found to be free of ectoparasites and clinical evidence of infectious or contagious diseases of equidae and, as far as can be determined, exposure thereto;.			
	II.14	it/they has\have not come into contact with any animals, products or equipment of a lesser zoosanitary health status during the entire required periods of residency, isolation, transportation to the port of exportation and loading onto the international transport carrier and the carrier has been instructed to maintain this status throughout transport to Canada			
N - 4	II.15	it/they has\have been treated before and at the time of loading in accordance with the relevant provisions of Regulation (EC) No 1/2005, in particular as regards watering and feeding and it/they are fit for the intended transport.			
Notes					
Part I:	Box no. I.11:	Indicate the premises of export and/or pre-export isolation facility, if different.			
	Box no. I.28:	Identification system: insert "Passport in accordance with Commission Regulation (EC) No 504/2008" or describe the other recognised (e.g FEI passport, breed registry, etc.) means of identification (which clearly and uniquely identifies the animal, and includes verifiable visual characteristics) used, and "microchip". Specify where the microchip is located.			
		Identification number: shall correspond to the alpha-numeric code of the microchip displayed by the appropriate reading device. If there is a unique number associated with the second means of identification (e.g. passport number), it should be recorded on the accompanying export health certificate.			
		According to the import rules of Canada, the animal must be marked with a microchip. The number of the microchip must be recorded on the accompanying export health certificate and, when possible, on the second means of identification.			
		For the verification of the identity of the animal it is mandatory to make available at the point of entry into Canada a reading device capable of reading and displaying the alpha- numeric code inserted in Box I.28, unless the microchip used is an ISO microchip.			
Part II:					
(1)	Delete as	appropriate.			
(2)	Geldings	and equidae 731 days of age or less are exempt from CEM testing.			
(3)	Check aga	ainst wording of corresponding Canadian Import Permit.			
(4)	nens must have been collected by a licensed veterinarian under the supervision of an official ian and were submitted in Amies transport medium with charcoal, transported refrigerated ozen, and cultured for CEM within 48 hours of collection in a laboratory officially approved to or CEM. During transport to the laboratory the specimens were accompanied by a statement the veterinarian collecting the specimens indicating the date and time of their collection.				
(5)		ine animal(s) has/have undergone any form of antibiotic treatment, collection of specimens for ng (swabs) must not commence until a minimum of seven (7) days post treatment.			
(6)	In the laboratory the specimens must be cultured for a minimum of 7 days (starting when the samples are cultured to laboratory media) on Eugon agar with 10% chocolated horse blood and onto the same medium with the following selective inhibitors: amphotericin-B (5μ g/ml), trimethoprim (1μ g/ml) and clindamycin (5μ g/ml). The plates must be incubated at 37°C in an atmosphere of 5 to 10 percent carbon dioxide and examined for gross contamination at 24 and 48 hours. The plates must be examined for suspect CEM organism colonies after 72 hours incubation and at 48-hour intervals thereafter. If no suspect colonies are observed after at least 168 hours of incubation, specimens should be reported as "CEM organism was not isolated.				
(7)	An officia	l copy of the laboratory report on CEM testing must be attached to this certificate			
. ,					
Certifying O	litter				

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