CHAPTER 4(A)

Health certificate

For the import of blood and blood products from equidae to be used outside the feed chain, for dispatch to or for transit through $^{(2)}$ the European Union

COUNTRY: UNITED STATES

Veterinary certificate to EU

	I.1. Consignor Name Address	I.2. Certificate reference No I.2.a.		
		I.3. Central competent authority APHIS-VS		
	Tel.	I.4. Local competent authority		
	1.5. Consignee Border inspection post through which consignment is intended to leave the EU	I.6. Person responsible for the load in EU Name Address		
Part I : Details of dispatched consignment	Name Address	Postal code Tel.		
	Postal code Tel.			
	I.7. Country of ISO code I.8. Region of Code origin US US-0	I.9. Country of ISO I.10. Region of eode destination code destination		
atche	I.11. Place of origin	I.12. Place of destination		
s of dispa	Name Approval number Address	Custom warehouse Name Approval number Address		
Detail		Postal code		
Part I:				
	Name Approval number Address			
	Name Approval number Address			
	I.13. Place of loading	I.14. Date of departure		
	I.15. Means of transport	I.16. Entry BIP in EU		
	Aeroplane □ Ship □ Railway wagon □ Road vehicle □ Other □	1.17.		
	Identification	1.17.		
	Documentation references			
	I.18. Description of commodity	I.19. Commodity code (HS code)		
		I.20. Quantity		
	I.21. Temperature of product	I.22. Number of packages		
	Ambient ☐ Chilled ☐ Frozen ☐			

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COUNTRY: UNITED STATES

Blood and blood products from equidae for purposes outside the feed chain

	I.2. Certificate reference No	I.2.a.
I.23. Seal/Container No	I.24. Type of packaging	
I.25. Commodities certified for:	_	
Technical use □		
I.26. For transit through EU to third country	I.27. For import or admission into EU	
Third country ISO code		
I.28. Identification of the commodities		
Species (Scientific name)	Approval number of establishments Manufacturing plant	

Page ____ of ____

II. Health information II.a. Certificate reference No II.b.

I, the undersigned official veterinarian, declare that I have read and understood Regulation (EC) No 1069/2009 of the European Parliament and of the Council^(1a) and in particular Article 8(c) and Article 8(d) and Article 10 thereof, and Commission Regulation (EU) No 142/2011^(1b), and in particular Chapter IV of Annex XIII thereto, and certify that the blood or blood products of equidae described above:

- II.1. consist of blood or blood products from equidae that satisfy the health requirements below;
- II.2. consist exclusively of blood or blood products of equidae not intended for human or animal consumption;
- II.3. have been obtained from animals that originate from the EU Member States or from a third country, territory or part thereof listed in the column 'third countries' lists' of row No 3 of Table 2 in Section 1 of Chapter II of Annex XIV to Regulation (EU) No 142/2011 where the following diseases are compulsorily notifiable: African horse sickness, dourine, glanders (*Burkholderia mallei*), equine encephalomyelitis (all types including Venezuelan equine encephalomyelitis), equine infectious anaemia, vesicular stomatitis, rabies, anthrax;
- II.4. have been derived from blood from equidae, which was collected under the supervision of a veterinarian in slaughterhouses approved in accordance with Regulation (EC) No 853/2004 of the European Parliament and of the Council⁽³⁾, in slaughterhouses approved and supervised by the competent authority of the country of collection and in facilities approved and supervised by the competent authority of the country of collection for the purpose of collecting blood from equidae for the production of blood products for purposes other than feeding for farmed animals;
- II.5. have been derived from blood which was collected from equidae:
- II.5.1. which on inspection on the date of blood collection did not show clinical signs of any of the compulsorily notifiable diseases listed in Annex I to Council Directive 2009/156/EC⁽⁴⁾, and of equine influenza, equine piroplasmosis, equine rhinopneumonitis and equine viral arteritis listed in point 4 of Article 1.2.3 of the Terrestrial Animal Health Code of the World Organisation for Animal Health (OIE), 2010 edition;
- II.5.2. which have been kept for at least 30 days prior to the date of and during blood collection on holdings under veterinary supervision which were not subject to a prohibition order pursuant to Article 4(5) or restrictions for African horse sickness in accordance with Article 5 of Directive 2009/156/EC;
- II.5.3. which had no contact with equidae from a holding which was subject to a prohibition order for animal health reasons pursuant to Article 4(5) of Directive 2009/156/EC;
- II.5.4. for which the period for the prohibition order referred to in points II.5.2. and II.5.3 has been determined as follows:

(2) either

[not all the animals of species susceptible to the disease located on the holding have been slaughtered, in which case the period of prohibition must be at least:

- six months in the case of glanders (*Burkholderia mallei*), beginning on the date on which the equidae infected with the disease are slaughtered,
- six months in the case of equine encephalomyelitis of any type, including Venezuelan equine encephalomyelitis, beginning on the date on which the equidae infected with the disease are slaughtered,
- in the case of equine infectious anaemia, until the date on which, the infected animals having been slaughtered, and the remaining animals have shown a negative reaction to two Coggins tests carried out three months apart,
- six months from the date of the last recorded case of vesicular stomatitis,
- one month from the date of the last recorded case of rabies,
- 15 days from the date of the last recorded case of anthrax;]

⁽²⁾or

[all the animals of species susceptible to the disease located on the holding have been slaughtered and the premises were disinfected, in which case the period of prohibition must be 30 days, beginning on the date on which the animals were slaughtered and the premises disinfected, except in the case of anthrax, where the period of prohibition shall be 15 days;]

- II.6. blood products come from an establishment or plant approved or registered by the competent authority of the third country meeting the specific conditions set out in Article 23 or 24 of Regulation (EC) No 1069/2009;
- II.7. blood products have been produced from blood which fulfils the conditions referred in II.4 and II.5 and

(2) either

[has been collected from equidae which have been kept for a period of at least three months, or since birth if less than three months old, prior to the date of collection on holdings under veterinary supervision in the country of collection which during that period and the period of blood collection has been free of:

- (a) African horse sickness for two years;
- (b) Venezuelan equine encephalomyelitis for a period of at least two years;
- (c) glanders
 - (2) either [for a period of three years;]

COUNTRY: UNITED STATES

Blood and blood products from equidae for purposes outside the feed chain

II.	Health info	ormation	II.a. Certificate reference No	II.b.	
		inspection for careful examin cavities and sin	is six months where the animals have passed of glanders in the slaughterhouse referred to in action of mucous membranes from the trache uses and their ramifications, after splitting the heing the nasal septum;	II.4, including a a, larynx, nasal	
		(d) in the case of blood production months;]]	ducts other than serum and plasma, vesicular	stomatitis for six	
	⁽²⁾ or	check, for the inactivation of po- encephalomyelitis of all types in anaemia, vesicular stomatitis and		sickness, equine equine infectious	
		(0)	at a temperature of 65°C for at least three hours;		
		-	kGy by gamma rays;]		
		(0)	pH 5 for two hours;]		
	- 11	•	of at least 80°C throughout their substance;]]		
II.8.		ions have been taken to avoid oing production, handling and packa	ontamination of the blood and blood products	with pathogenic	
II.9.	blood and blood products were packed in sealed impermeable containers clearly labelled 'NOT FOR HUMAN OR ANIMAL CONSUMPTION' and bearing:				
		•	er of the establishment of collection;		
	(b) in the	e case of blood products, the appro	val number of the establishment of production;		
II.10.	the product	ts were stored in enclosed storage			
Notes					
Part I:	Box reference I.6: Pers		propean Union: this box is to be filled in only if it is a certificate for	or transit commodity; it	
_	•	ertificate is for import commodity. I I.12: Approval number: the registration numb	er of the establishment or plant, which has been issued by the co	mpetent authority.	
-	Box reference I.12: Pla		y if it is a certificate for transit commodity. The products in transi		
-	Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor must inform the BIP of entry into the EU.				
		opriate Harmonized System (HS) code under to	he following heading: 30.02. e seal number (if applicable) must be included.		
-		hnical use: any use other than for animal cons			
-		I.27: fill in according to whether it is a transit	or an import certificate.		
-	Box reference I.28: (a) Manufacturing p	alant.			
		ase of blood, provide the approval number of t	he registered establishment of collection.;		
	. ,	ase of blood products, provide the approval nu	•		
Part II:		amongst the following: Equus cabalus, Equus	asınus, Equus cabalusXasınus.		
(1a)	OJ L 300, 14.11.2009,	p. 1.			
(1b) (2)	OJ L 54, 26.2.2011, p.				
(3)	Delete as appropriate. OJ L 139, 30.4.2004, p				
(4)	OJ L 192, 23.7.2010, p				
	-	tamp must be in a different colour to that of the			
		sponsible for the consignment in the Europe thes the border inspection post.	ean Union: this certificate is only for veterinary purposes and	must accompany the	
Offici	al veterinarian/Offi	icial inspector			
	Name (in capital	letters): Quali	ication and title:		
	(sapital	gaan			
	Date:	Sign	ature:		
	Date.	-			
		Starr	ıp:		