### CHAPTER 4 (D)

#### **Health certificate**

For treated blood products, excluding those of equidae, for the manufacture of technical products, intended for dispatch to or for transit through<sup>(2)</sup> the European Union

CO	UNTRY			Veterinary certificate to EU	
	I.1. Consignor  Name  Address		I.2. Certificate reference number	I.2.a	
<del> </del>			I.3. Central Competent Authority		
Part I : Details of dispatched consignment			1.4. Local Competent Authority		
ign	Tel.N°				
ons	I.5. Consignee Name		1.6. Person responsible for the consignment in EU     Name		
q c					
che	Address  Postal code  Tel.N°		Address Postal code		
pat			Tel.N°		
disp	I.7.Country of origin ISO code I.8. Reg	ion of origin Code	I.9. Country of destination ISO	code I.10.	
of	I.11. Place of origin	<u> </u>	I.12. Place of destination		
ails					
Det			Custom warehouse	Custom warehouse	
		pproval number	Name	Approval number	
art	Address		Address		
P			Postal code		
	I.13. Place of loading		I.14. Date of departure		
	I.15. Means of transport		I.16. Entry BIP in EU		
	Aeroplane Ship Road vehicle	Railway wagon Other			
	Identification:		I.17.		
	Documentary references:  I.18. Description of commodity		1.19. Commodity code (HS code) 30.02		
	1.16. Description of commonty				
				I.20.Quantity	
	I.21 Temperature of product			I.22. Number of packages	
	Ambient	Chilled	Frozen	I.22. Number of packages	
		Chilled	Frozen		
	Ambient	Chilled	Frozen	I.22. Number of packages	
	Ambient	Chilled Technical use	Frozen	I.22. Number of packages	
	Ambient		Frozen	I.22. Number of packages	
	Ambient		Frozen	I.22. Number of packages	
	Ambient   1.23. Identification of container/Seal number  1.25. Commodities certified for:  1.26. For transit to 3rd Country vis-à-vis EU	Technical use		I.22. Number of packages	
	Ambient			I.22. Number of packages	
	Ambient   1.23. Identification of container/Seal number  1.25. Commodities certified for:  1.26. For transit to 3rd Country vis-à-vis EU	Technical use		I.22. Number of packages	
	Ambient	Technical use	1.27. For import or admission into EU	I.22. Number of packages	
	Ambient	Technical use		I.22. Number of packages	
	Ambient	Technical use	1.27. For import or admission into EU  Approval number of establishments	I.22. Number of packages  I.24. Type of packaging	
	Ambient	Technical use	1.27. For import or admission into EU  Approval number of establishments	I.22. Number of packages  I.24. Type of packaging	
	Ambient	Technical use	1.27. For import or admission into EU  Approval number of establishments	I.22. Number of packages  I.24. Type of packaging	
	Ambient	Technical use	1.27. For import or admission into EU  Approval number of establishments	I.22. Number of packages  I.24. Type of packaging	
	Ambient	Technical use	1.27. For import or admission into EU  Approval number of establishments	I.22. Number of packages  I.24. Type of packaging	
	Ambient	Technical use	1.27. For import or admission into EU  Approval number of establishments	I.22. Number of packages  I.24. Type of packaging	

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# $\label{thm:conditional} \textbf{Treated blood products, excluding those of equidae, for technical products}$

II.	Health information	II.a.	Certificate reference number:	II.b.				
	I, the undersigned official veterinarian, declare that I have read and understood Regulation (EC) No1774/2002 <sup>(1)</sup> and in particular Article 4(1)(c), Article 6 and Chapter IV of Annex VIII thereof and certify that:							
II.1.	the blood products described above consist of blood products that satisfy the requirements below;							
II.2.	they consist exclusively of blood products not intended for human or animal consumption;							
II.3.	they have been prepared and stored in a plant approved, validated and supervised by the competent authority in accordance with Article 18 or in the establishment of collection and where appropriate Article 11 of Regulation (EC) No 1774/2002 <sup>(2)</sup> , exclusively with the following animal by-products:							
<sup>(2)</sup> either	[- blood of slaughtered animals, which is fit for human consumption in accordance with Union legislation, but is not intended for human consumption for commercial reasons;]							
<sup>(2)</sup> and/or	[- blood of slaughtered animals, which is rejected as unfit for human consumption but is not affected by any signs of diseases communicable to humans or animals, derived from carcasses that are fit for human consumption in accordance with Union legislation;]							
<sup>(2)</sup> and/or	[- blood obtained from animals other than ruminants that are slaughtered in a slaughterhouse, after undergoing ante-mortem inspection, and were fit, as a result of such inspection, for slaughter for human consumption in accordance with Union legislation;]							
<sup>(2)</sup> and/or								
II.4.	the blood from which such products are manuf							
<sup>(2)</sup> either	[in slaughterhouses approved in accordance w	ith Union le	egislation,]					
(2)or	[in slaughterhouses approved and supervised b							
$^{(2)}or$	[from live animals in facilities approved and su	•						
<sup>(2)</sup> [II.5.	In case of blood products derived from <i>taxa Artiodactyla</i> , <i>Perissodactyla</i> and <i>Proboscidea</i> including their crossbreeds, other than <i>Suidae</i> and <i>Tayassuidae</i> , the products have undergone one of the following treatments, guaranteeing the absence of pathogens of foot-and-mouth disease, vesicular stomatitis, rinderpest, peste des petits ruminants, Rift Valley fever and bluetongue:							
<sup>(2)</sup> either	[heat treatment at a temperature of 65°C for at	least three	hours, followed by an effectiveness che	eck,]				
$^{(2)}or$	[irradiation at 25 kGy by gamma rays, followed by an effectiveness check,]							
$^{(2)}or$	[change in pH to pH 5 for two hours, followed	by an effec	ctiveness check,]					
$^{(2)}or$	[heat treatment of at least 80°C throughout the	ir substanc	e, followed by an effectiveness check]]					
<sup>(2)</sup> [II.6.	In the case of blood products derived from <i>Suidae</i> , <i>Tayassuidae</i> , poultry and other avian species, the products have undergone one of the following treatments guaranteeing the absence of pathogens of the following diseases: foot-and-mouth disease, vesicular stomatitis, swine vesicular disease, classical swine fever, African swine fever, Newcastle disease and highly pathogenic avian influenza as appropriate to the species;							
<sup>(2)</sup> either	-	least three hours, followed by an effectiveness check,]						
$^{(2)}or$	[irradiation at 25 kGy by gamma rays, followed by an effectiveness check,]							
<sup>(2)</sup> or	[heat treatment of at least 80°C for <i>Suidae/Tayassuidae</i> <sup>(2)</sup> and at least 70°C for poultry and other avian species <sup>(2)</sup> throughout their substance, followed by an effectiveness check]]							
<sup>(2)</sup> [II.7.	In the case of blood products derived from species other than listed under II.5. or II.6. the products have undergone of the following treatment (please specify):;]							
II.8.	the products were:							
(2)either	[packed in new or sterilised bags or bottles,]							
<sup>(2)</sup> or	[transported in bulk in containers or other means of transport that were thoroughly cleaned and disinfected with a disinfectant approved by the competent authority before use,]							
	the outer packaging or containers bear labels in	ndicating "l	NOT FOR HUMAN OR ANIMAL CO	NSUMPTION";				
II.9.	the products were stored in enclosed storage;							
II.10.	the products have undergone all precautions to	avoid cont	amination with pathogenic agents after	treatment.				
Notes								
Part I:								
•	Box reference I.6: Person responsible for the	consignmen	nt in the European Union: this box is to	be filled in only if it				

#### COUNTRY

# $\label{thm:conditional} \textbf{Treated blood products, excluding those of equidae, for technical products}$

II.	Health information	II.a.	Certificate reference number:	II.b.				
	is a certificate for transit commodity; it may be	filled in	if the certificate is for import commodity	у.				
•	Box reference I.12: Place of destination: this box is to be filled in only if it is a certificate for transit commodity. The products in transit can only be stored in free zones, free warehouses and custom warehouses approved for that purpose.							
•	Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship); information is to be provided in case of unloading and reloading.							
•	Box reference I.23: for bulk containers, the container number and the seal number (if applicable) must be included.							
•	Box reference I.26 and I.27: fill in according to	Box reference I.26 and I.27: fill in according to whether it is a transit or an import certificate.						
Part	П:							
(1)	OJ L 273, 10.10.2002, p. 1.							
(2)	Delete as appropriate.							
•	The signature and the stamp must be in a differ	The signature and the stamp must be in a different colour to that of the printing.						
•	Note for the person responsible for the consignment in the European Union: this certificate is only for veterinary purposes and must accompany the consignment until it reaches the border inspection post.							
Offic	cial veterinarian							
	Name (in capital letters):		Qualification and title:					
	Date:		Signature:					
	Stamp:							