## Komunikat Głównego Lekarza Weterynarii w sprawie środków ochronnych dotyczących wymagań zdrowia zwierząt podjętych przez Komisję europejską celem zabezpieczenia Unii Europejskiej przed przeniknięciem na jej terytorium Epidemicznej Biegunki Świń (PDE)

Epidemiczna biegunka świń (PDE) jest chorobą powodowaną przez deltakoronawirus występujący w Azji i Ameryce Północnej. Choroba ta nie była notowana w Unii Europejskiej.

Suszone produkty z krwi świńskiej importowane z krajów trzecich mogą być wykorzystywane jako materiał paszowy w żywieniu prosiąt i trzody chlewnej.

Mając na uwadze powyższe, oraz niebezpieczeństwo przeniesienia wirusa PDE na trzodę chlewną w UE, Komisja Europejska ustaliła specjalne warunki jakie muszą zostać spełnione przez produkty z krwi pochodzącej od świń, importowane do UE z przeznaczeniem do produkcji pasz dla prosiąt i trzody chlewnej, obejmujące specjalną obróbkę surowca.

Informuję, że publikacja rozporządzenia KE wprowadzającego wspomniane środki ochronne oraz wzór świadectwa zdrowia dla produktów z krwi pozyskanej od świń, nieprzeznaczonej do spożycia przez ludzi, które mogą być wykorzystane jako materiał paszowy przewidywana jest na 13 maja br.

W związku z powyższym wszystkie przesyłki wspomnianych produktów certyfikowane po dniu publikacji w/w rozporządzenia muszą być zaopatrzone w nowy wzór świadectwa i spełniać określone w nim specjalne warunki!

Poniżej prezentuję projekt nowego świadectwa zdrowia

Główny Lekarz Weterynarii

#### **ANNEX**

#### **Health certificate**

For blood products not intended for human consumption that could be used as feed material, intended for dispatch to or for transit through<sup>2</sup> the European Union

CO	UNTRY:	Veterinary certificate to EU			
Part I: Details of dispatched consignment	I.1. Consignor Name	I.2. Certificate reference No	I.2.a.		
	Address	I.3. Central competent authority			
	Tel.	I.4. Local competent authority			
	I.5. Consignee Name Address  Postcode Tel.	I.6. Person responsible for the load in El Name Address Postcode Tel.	U		
	I.7. Country of ISO code I.8. Region of origin Code origin	I.9. Country of destination ISO code	I.10. Region of Code destination		
ails	I.11. Place of origin	I.12. Place of destination			
art I : Deta	Name Approval number Address Name Approval number Address		Custom warehouse  Approval number		
Ь	Name Approval number	Postcode			
	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 ☐ Identification  Documentation references	I.17.			
•	I.18. Description of commodity	I.19. Commodity code (HS code)			
			I.20. Quantity		
	I.21. Temperature of product Ambient □ Chilled □	Frozen	I.22. Number of packages		
,	I.23. Seal/Container No		I.24. Type of packaging		
i	I.25. Commodities certified for:				
	Animal feedingstuff □ Techn	nical use			
	I.26. For transit through EU to third country	I.27. For import or admission into E	U		
	Third country ISO code				
	I.28. Identification of the commodities	l			
	Species (Scientific name) Nature of commodity	Approval number of establishments Manufacturing plant	Batch number		

**COUNTRY** 

# Part II: Certification

II.	Health info	ormation II.a. Certificate reference No II.b.			
II.1. II.2. II.3.	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 Commission Regulation (EU) No 142/2011 (1b) and certify that the blood products described above: consist of blood products that satisfy the health requirements below; consist exclusively of blood products not intended for human consumption; have been prepared and stored in a plant, approved, validated and supervised by the competent authority in accordance with Article 24 of Regulation (EC) No 1069/2009; have been prepared exclusively with the following animal by-products:				
	(2) 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 in accordance with Union legislation, but which did not show any signs of diseases communicable to humans or animals, derived from carcases that have been slaughtered in a slaughterhouse and were considered fit for human consumption following an ante-mortem inspection in accordance with Union legislation;]			
II.5.	in order to i	nactivate pathogenic agents, have been submitted			
	<sup>(2)</sup> either	[to processing in accordance with processing method			
	<sup>(2)</sup> or	[to a method and parameters which ensure that the product complies with the microbiological standards set in Chapter I of Annex X to Regulation (EU) No 142/2011;]			
	<sup>(2)</sup> or	[in the case of blood products, including spray dried blood and blood plasma, of porcine origin intended for the feeding of porcine animals, to a heat treatment at a temperature of at least 80°C throughout the substance and the dry blood and blood plasma is of not more than 8% moisture with a water activity (Aw) of less than 0,60.]			
II.6.	have been examined under the responsibility of the competent authority taking a randon sample immediately prior to dispatch and found it to comply with the following standards <sup>(4)</sup> :				
	Salmonella	absence in 25g: $n = 5$ , $c = 0$ , $m = 0$ , $M = 0$ ,			
	Enterobacteriaceae: $n = 5$ , $c = 2$ , $m = 10$ , $M = 300$ in 1 gram;				
II.7. the end product was:		duct was:			
	<sup>(2)</sup> either	[packed in new or sterilised bags;]			
	<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,]			
		pear labels indicating 'NOT FOR HUMAN CONSUMPTION';			
II.8.	-	duct was stored in enclosed storage;			
II.9.	the product has undergone all precautions to avoid contamination with pathogenic agents after treatment;				
	<sup>(2)</sup> and	[in the case of spray dried blood and blood plasma of porcine origin intended for the feeding of porcine animals, has been stored in dry warehouse conditions under room temperature for at least 6 weeks.]			
II.10.		ntain and is not derived from:			
	<sup>(2)</sup> either	[specified risk material or mechanically separated meat obtained from bones of bovine, ovine or caprine animals and, except for animals born, continuously reared and slaughtered in a country or region classified as posing a negligible BSE risk by a decision in accordance with Article 5(2) of Regulation (EC) No 999/2001 of the European Parliament and of the Council (5), the animals from			

#### **COUNTRY**

### Blood products not intended for human consumption that could be used as feed material

II.	Health info	rmation	II.a. Certificate reference No	II.b.
	<sup>(2)</sup> or	slaughtered after stu killed by the same tissue by means of cranial cavity.] [bovine, ovine and of born, continuously re	by-product or derived product is donning by means of gas injected into method or slaughtered by laceration elongated rod-shaped instrume caprine materials other than those eared and slaughtered in a country as 3SE risk by a decision in accordance 1999/2001.]	o the cranial cavity or on of central nervous nt introduced into the derived from animals or region classified as

#### Notes

#### Part I:

- Box reference I.6: Person responsible for the consignment in the European Union: this box is to be filled in only if it is a certificate for transit commodity; it may be filled in if the certificate is for import commodity.
- 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.
- 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.19: use the appropriate HS code: 05.11.91 or 05.11.99.
- Box reference I.23: for bulk containers, the container number and the seal number (if applicable) should be included.
- Box reference I.25: technical use: any use other than for animal consumption.
- Box reference I.26 and I.27: fill in according to whether it is a transit or an import certificate.
- Box reference I.28: Species: select from the following: Aves, Ruminantia, Suidae, Mammalia other than Ruminantia, Pesca, Reptilia.

#### Part II:

- <sup>(1a)</sup> OJ L 300, 14.11.2009, p. 1.
- <sup>(1b)</sup> OJ L 54, 26.2.2011, p. 1.
- (2) Delete as appropriate.
- (3) Insert method 1 to 5 or 7 as applicable.
- (4) Where:
  - n = number of samples to be tested;
  - m = threshold value for the number of bacteria; the result is considered satisfactory if the number of bacteria in all samples does not exceed m;
  - M = maximum value for the number of bacteria; the result is considered unsatisfactory if the number of bacteria in one or more samples is M or more; and
  - c = number of samples the bacterial count of which may be between m and M, the sample still being considered acceptable if the bacterial count of the other samples is m or less.
- <sup>(5)</sup> OJ L 147, 31.5.2001, p. 1.
- 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 has to accompany the consignment until it reaches the border inspection post.

#### **COUNTRY**

## Blood products not intended for human consumption that could be used as feed material

II.	Health information	II.a. Certificate refer	rence No	II.b.
Official veterinarian/Official inspector				
Name (in capital letters):		Qualification and title:		
Date:		Signature:		
S	tamp:			