



Part I : Details of dispatched consignment

I.1. Consignor Name Address Country		I.2. Certificate reference number GBR	
		I.3. Central Competent Authority	
		I.4. Local Competent Authority	
I.5. Consignee Name Address Country		I.6. No.(s) of related original certificates	No.(s) of accompanying documents
I.7. Country of origin	ISO code	I.8. Region of origin	
		I.9. Country of destination	ISO code
		I.10. Region of destination	
I.11. Place of origin Name Address Country		I.12. Place of destination	
I.13. Place of loading		I.14. Date and time of departure	
I.15. Means of transport Aeroplane <input type="checkbox"/> Ship <input type="checkbox"/> Railway wagon <input type="checkbox"/> Road vehicle <input type="checkbox"/> Other <input type="checkbox"/>		I.16. Entry Point	
Identification: Number(s):		I.17. CITES	
I.18. Temperature of products		I.19. Total Gross Weight	I.20. Total number of packages
I.21. Seal/Container number			
I.22. Commodities certified for : Animal Feeding stuff <input type="checkbox"/> Technical Use <input type="checkbox"/> Other end-use <input type="checkbox"/>			
I.23. Transit through 3rd country		I.24. For Export <input type="checkbox"/>	
I.25. Identification of the commodities Customs Code Title			
Species (Scientific name)	Approval No of Establishment	Description of Product	Date of Production



Part II: Certification

II. Health information

II.a. Certificate reference number

GBR

I, the undersigned official veterinarian, after due inquiry and to the best of my knowledge, do hereby certify that the processed porcine blood described within this certificate meets all the conditions laid down in the health attestations below:

- II.1. The country or zone of origin is free of foot-and-mouth disease, classical swine fever, swine vesicular disease, and African swine fever. Vaccination against these diseases is prohibited in that country.
- II.2. The products were only produced and exported to Canada by the producer and exporter identified in the Import Permit and mentioned on this certificate's first page.
- II.3. The certified rendered products were produced:
 - (2) either [II.3.1. In a dedicated facility that does not receive, process or store any ruminants and things derived from ruminants, and the product has been prepared, processed, packaged, stored, shipped and otherwise handled in a manner to avoid contamination with any ruminant tissues or things derived from ruminants,]
 - (2) or [II.3.1. on a dedicated line from receipt of raw material to final packaging and storage with no risk of cross contamination with any ruminants and things derived from ruminants, and the product has been prepared, processed, packaged, stored, shipped and otherwise handled in a manner to avoid contamination with any ruminant tissues or things derived from ruminants (if the facility has ineligible materials on the premises, all rendered products for export to Canada must be produced on a totally dedicated line.)]
- II.4. The blood used in the certified products was obtained only from carcasses that passed ante-mortem inspection and were subjected to post-mortem inspection in slaughterhouses approved and supervised by the competent authority.
- II.5. None of the animals from which any of the unprocessed blood used to manufacture the rendered products are derived from premises that were under any official restrictions by the UK and/ or EU Member State competent veterinary authority for any serious epizootic disease to which the species from which the product or by-product was derived is susceptible and can be transmitted by the untreated product or by-product and none of the animals from which the animal origin raw materials are derived were under movement restrictions for or were culled or eradicated as part of a disease response for any reportable disease as defined by Canada (1).
- II.6. The raw materials used to produce the rendered products have been transported in dedicated vehicles.
- II.7. The porcine blood products have been processed in an establishment approved by the veterinary competent authority of the United Kingdom for export to Canada.
- II.8. The blood products are exclusively of porcine origin.
- II.9. The blood products have been heat treated to at least 80° Celsius throughout their substance.
- II.10. Each shipment to Canada has been tested negative for the presence of ruminant DNA with a PCR method, in an accredited laboratory.
- II.11. The porcine blood products have been packed using new and secure packaging materials.
- II.12. The raw materials and additives used in the porcine blood products have been clearly indicated on the exterior packaging.
- II.13. The porcine blood products for export have been produced, processed, stored and transported in such a manner as to prevent contamination by communicable animal disease pathogens up to the point of departure from the country of origin.
- II.14. The product label bears the following statement "The product does NOT contain prohibited material, as defined by section 162 of the Canadian Health of Animals Regulations².

Notes

All pages must be presented at least in English and/or French. The official stamp and the signature of the official veterinarian must appear on each separate sheet, including any attached lists. The signature and the stamp must be in a different color to that of the printing.

Part I:

- Box reference I.1: Indicate the details of the exporter.
 - Box reference I.2.: Indicate the unique reference number.
 - Box reference I.5.: Indicate the details of the importer.
 - Box reference I.6.: Indicate the CFIA Import permit number
 - Box reference I.11.: Place of origin: name and address of the dispatch establishment.
 - Box reference I.15.: Indicate the names of the ships and, if known, the flight numbers of aircraft.
 - Box reference I.19.: Indicate total gross weight and total net weight.
 - Box reference I.21.: For containers or boxes, the container number and the seal number should be included.
 - Box reference I.22.: Commodities certified for must identify the end use.
 - Box reference I.25.: Customs code and title: Use the appropriate Harmonized System (HS) code under the following headings: 0511, 3002, 3502, 2301
- Processing plant: Indicate the establishment approval number.
- Description of product: as per Import permit
- Animal species: only porcine blood allowed



II. Health information

II.a. Certificate reference number

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Date of production: Shall be indicated in the following format: dd/mm/yyyy

Part II:

- (1) The CFIA list of diseases that are reportable in Canada can be found on the CFIA website: Animal Health Status By Disease - Animals - Canadian Food Inspection Agency. The CFIA accepts the OIE classification of BSE Risk Status: List of BSE risk status: OIE - World Organisation for Animal Health.
- (2) Where the product is destined for use in livestock feed, it is approved for use and listed in Schedule IV of the Feeds Regulations and is labelled as required by the Feed Regulations, which have been captured here: https://ec.europa.eu/food/safety/international_affairs/agreements/export-library_en

Part II: Certification

8609EHC APPLICATION

Official veterinarian or official inspector

Name (in Capital):
Local Veterinary Unit:
Date:
Stamp

Qualification and title:
LVU N°:
Signature: