

# **DRAFT VETERINARY HEALTH CERTIFICATE FOR IMPORT<sup>1</sup> OF BOVINE SERUM INTO INDIA**

## **I. General Information**

1. Veterinary Health certificate No: Date:		2. Competent Authority 2.1 Ministry: 2.2 Department: 2.3 Contact Details and Email:	
3. Invoice No. Date:		4. Quantity: 4.1 Type of packaging 4.1 Number of packages 4.3 Net Weight	
5. Consignor / exporter Name: Address: Tel. no. and Email:			
6. Consignee /importer: Name: Address: Tel. no. and Email:			
7. Country of origin:		ISO Code:	
8. Place of loading:			
9. Country of Destination:		ISO Code:	
10. Declared Port of Entry <sup>2</sup>			
11. Mode of Transport:		12. Temperature:	
13. Identification of Container:		14. Seal No.	
15. SIP/DGFT License no. with Date and Validity			
16. Identification of the product as described below: (multiple lines may be used for multiple products)			
a)	Description of the Products along with HS code:	Intended purpose:	
b)	Name and address of Manufacturer / Establishment	Approval number/s of establishment /s (Number /Date / Validity) along with Name and address of the Registration / Accreditation Authority	
c)	Name of the product	Lot no. / Batch no.	
d)	Date of Manufacture or Packaging	Best before (if applicable)	Date of Expiry

<sup>1</sup> Import of livestock products into India is subjected to fulfillment of the Live-stock Importation Act, 1898 and the rules / regulations there under as notified time to time.

<sup>2</sup>Port of Entry as notified by Ministry of Fisheries, Animal Husbandry and Dairying, Government of India considering applicability of Sanitary Import Permit (SIP) or not, as the case may be.

## II. Sanitary information

The undersigned Official Veterinarian certifies that the products described above satisfy the following requirements:

- i. The country of export/origin is under negligible Bovine Spongiform Encephalopathy (BSE) risk in accordance with Chapter 11.4. of the Terrestrial Code.  
or  
Bovine serum has been obtained from the donor animals not subjected to a stunning process with a device injecting compressed air or gas into the cranial cavity, or to a pithing process, or to any other procedure that can contaminate the blood with nervous tissue, prior to slaughter and collected and processed in a manner that ensures they are not contaminated with nervous tissue.
- ii. The country of export/origin is free from BVD and Foot and Mouth Disease Virus as per WOAHP listing.  
or  
In case the import is from the BVD and Foot and Mouth Disease Virus positive country, the consignment should be accompanied with a certificate of quality analysis and tested negative for BVD and Foot and Mouth Disease Virus.
- iii. The product originates from donor animal which had undergone ante- and post-mortem inspections, which have been slaughtered for human consumption at an abattoir approved by the National Veterinary Authority of the country of origin in accordance with WOAHP Chapter 6.3 with favourable results and shows no clinical sign of any infectious and contagious diseases related to the source animal species including Anthrax, Foot and Mouth Disease, Lumpy Skin Disease.  
Or  
The product is obtained from the live and officially certified healthy donor animal originated from the officially approved establishments, which did not show any clinical sign and symptoms of any infectious/contagious diseases related to the source animal species including Anthrax, Foot and Mouth Disease, Lumpy Skin Disease.
- iv. The donor animals originate from an establishment at least six month prior to serum collection accredited by the exporting country with donor animals individually identified by a unique number of alphanumeric code, permanently applied to the animals by means of identification or tattoo, correlating with the serum collection documents and the identification numbers should be stated in this certificate.
- v. The product came from the donor animal which has not been kept in the infected zone and establishment for the diseases related to source animal

species and have not been derived from the source animal species slaughtered/culled as part of an eradication/control program for any disease related to source animal species.

- vi. The product has been collected by the expert and trained team approved by the competent authority of exporting country as per standard and approved procedures as approved by the official authority of exporting country as per international guidelines related to Laboratory bio-safety and bio-security
- vii. The product is for research purpose only and should not contain any infectious agent. Care has been taken to ensure that the product is produced and handled in a manner which ensures that the product do not contain and is not contaminated with any pathogenic agents or organism with no threat to outside environment and related species.
- viii. Guidelines for International transfer (IATA guidelines) including packaging (Triple layered) and laboratory containment of animal pathogenic agents shall also be followed as per WOAHP chapter 5.8 and spillage and leakage must be strictly avoided.
- ix. Adequate precautions were taken after collection, processing and during transit to avoid contact and contamination of product with any potential source of infection. The product is packed in new, fresh, clean packing material and the materials are not exposed to any products with potential source of infectious materials.
- x. Serum must be sterile and would be used for the intended purpose only.

Official stamp:

Signature\_\_\_\_\_

<b>Official Veterinarian</b>	
Name:	Designation:
Address and Email:	
Date:	

**Post import clearance requirements:**

1. On arrival in India the consignment and the documents will be examined by the Animal Quarantine and Certification Services. Department of Animal Husbandry and Dairying, Government of India.

2. In case of the documents are not conforming to the requirements or the product is not as per protocol, appropriate action shall be taken by the Department of Animal Husbandry and Dairying, Ministry of Fisheries, Animal Husbandry and Dairying, Government of India.
3. In case the import is from the BVD and Foot and Mouth Disease Virus positive country, the samples shall be taken for testing of BVD by ICAR-NIHSAD and FMD by ICAR-NIFMD respectively.
4. The material should be handled as per guidelines related to Laboratory biosafety and biosecurity guidelines and the destination laboratory shall have the regulatory approvals and proper biosafety levels.
5. The disposal if any shall be as per Bio-Medical Waste Management Rules.

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Date: 18<sup>th</sup> August, 2025