



#### **CERTIFICATE OF A PHARMACEUTICAL PRODUCT^1**

| No of Certificate : 3237540/TS/2022  | Valid UpTo: <b>09/08/2025</b>  |
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| 1.Name and Dosage form of Product:  Rabies Antiserum 1000 I.U. I.H.S. (5 mL Liquid Vial)   | Exporting(Certifying)Country: INDIA  |
| 1.1 Active Ingredients(s)^2 and amount(s) per unit dose^3:  Each mL contains Enzyme Refined; Equine Antirabies immunoglobulin fragments not less than 200 I.U. | Importing(Requesting) country <b>Belarus</b>   |
| For complete qualitative comp <mark>osition inc</mark> luding excipients see above^4   |  |
| 1.2 Is this Product licensed to be placed on the market for use in Exporting country   | ry?^5 Yes  |
| 1.3 Is this product actually on the marketing in the Exporting Country?  | Yes  |
| If the answer to 1.2 is Yes, continue with section 2A and omit section 2B.If the ans   | swer to 1.2 is No, omit section 2A continue with section   |
| 2A.1 Number of Product Licence^7: <b>01/MN/AP/2003/Sera/G, Dt. 17.12.2003,</b> valid up to 16.12.2023  | 2B.1 Applicant for Certificate(Name and Address)   |
| 2A.2 Product License Holder(Name and address): VINS Bioproducts Limited, Sy. No. 117, Thimmapur (V), Kothur(M), Ranga Reddy(Dist.), - 509325, Telangana, India | 2B.2 Status of Applicant^8   |
| 2A.3 Status of License Holders^8 : a   | 2B2.1 For Categories (b) and (c) the name and address of the manufacturer producing the dosage form is^9 |
| 2A.3.1 For Category b and c the name and address of the manufacturer producing the dosage form is ^9 NA  | 2B.3 Why is marketing authorisation lacking?   |
| 2A <mark>.4 Is</mark> Summary basis of approval appended ? ^10 Yes   | 2B.4 Remarks^13:   |
| 2A.5 Is the Attached, officially approved production information complete and consonant with the license?^11: Yes  |  |
| 2A.6 Applicant for Certificate, if different from licence holder (name and address) NA   |  |
| 3 Does t <mark>he ce</mark> rtifying authority arrange for periodic spection of the manufacturing form is produced?^14   |  |
| 3.1 Periodicity of routine inspection(years)   | Onco in a Year   |
| 3.2 Has the Manufacture of this type of dosage from been inspected?  | Yes  |
| 3.3 Do the facilities and operations conform to GMP as recommended by World I  | Health Organization?^15 Yes  |
| 4.0 Does the information Submitted by the applicant satisfy the certifying authorimanufacture of the product?^16   | ty on all aspects of the Yes   |
| Address of Certifying Authority.   | Name of the authorized person:   |
| Drugs Control Administration, Vengalraonagar, Hyderabad 500038, India Telephone No: 91-040-23814119 Fax No: 91-040-23814360                                    | Digitally Signed By  RAMDHAN GUGULOTH  |
|  | Deputy Director and Certifying Authority DRUGS CONTROL ADMINISTRATION                                    |

Date:19-08-2022 11:32:33 AM

**TELANGANA STATE** 





This certificate conforms to the format recommended by the World Health Organization (General instructions and explanatory notes overleaf)





#### **Explanatory notes**

- 1. This certificate, which is in the format recommended by WHO, establishes the status of the pharmaceutical product and of the applicant for the certificate in the exporting country. It is for a single product only since manufacturing arrangements and approved information for different dosage forms and different strengths can vary.
- 2. Use whenever possible, International Non-proprietary Names (INNs) or national non-proprietary names.
- 3. The formula (complete composition) of dosage form should be given on the certificate or be appended.
- 4. Details of quantitative composition are preferred, but their provision is subject to the agreement of the product licence holder.
- 5. When applicable, append details of any restriction applied to the sale, distribution or administration of the product that is specified in the product license.
- 6. Sections 2A and 2B are mutually exclusive.
- 7. Indicate when applicable, if the license is provisional, or the product has not yet been approved.
- 8. Specify whether the person responsible for placing the product on the market:
  - a) manufactures the dosage form;
  - (b) packages and/or labels a dosage form manufactured by an independent company; or
  - (c) is involved in non of the above
- 9.This information can be provided only with the consent of the product license holder or, in the case of non registered products, the applicant. Non-completion of this section indicates that the party concerned has not agreed to inclusion of this information. It should be noted that information concerning the site of production is part of the product license. If the production site is changed, the license Must be updated or it will cease to be valid.
  10. This refers to the document, prepared by some national regulatory authorities, that summarizes the technical basis on which the
- 10.This refers to the document, prepared by some national regulatory authorities, that summarizes the technical basis on which the product has been licensed.
- 11. This refers to the product information approved by the competent national regulatory authority, such as a Summary of Product Characteristics (SmPC).
- 12.In this circumstance, permission for issuing the certificate is required from the product license holder. This permission must be provided to the authority by the applicant.
- 13.Please indicate the reason that the applicant has provided for not requesting registration:
  - (a) the product has been developed exclusively for the treatment of conditions particularly tropical diseases not endemic in the country of export;
  - (b) the product has been reformulated with a view to improving its stability underP tropical conditions;
  - (c) the product has been reformulated to exclude excipients not approved for used in pharmaceutical products in the country of import;
  - (d) the product has been reformulated to meet a different maximum dosage limit for an active ingredient;
  - (e) any reason, please specify.
- 14. Not applicable means that the manufacture is taking place in a country other than that issuing the product certificate and inspection is conducted under the aegis of the country of manufacture.
- 15. The requirements for good practices in the manufacture and quality control of drugs referred to in the certificate are those included in the thirty
  - second report of the Expert Committee on Specifications for Pharmaceutical Preparations (WHO Technical Report Series No. 823, 1992

    Annex 1). Recommendations specifically applicable to biological products have been formulated by the WHO Expert Committee on Biological Standardization (WHO Technical Report Series, No. 822, 1992 Annex 1)
- 16.This section is to be completed when the product license holder or applicant conforms to status (b) or (c) as described in note 8 above. It is of Particular importance when foreign contractors are involved in the manufacture of the product. In these circumstances the applicant should supply the certifying authority with information to identify the contracting parties responsible for each stage of manufacture of the finished dosage form, and the extent and nature of any controls exercised over each of these parties.





