

# CERTIFICATE OF PHARMACEUTICAL PRODUCTS

No. of Certificate : HFW-H (DRUGS) 427/05/21-47      Exporting (certifying) Country : INDIA  
Valid up to : 22.02.2024      Importing (requesting) Country : PHILIPPINES

1.0 Proprietary Name (If applicable) and Dosages form of Product : PHRINTEC  
Phenylephrine Hydrochloride Injection USP 10mg/ml

Active ingredients(s) and amount per unit dose : Each ml contains:  
Phenylephrine Hydrochloride USP ..... 10 mg  
Sodium Metabisulphite USP.....2 mg  
Trisodium Citrate Dihydrate USP.....4 mg  
Citric acid Monohydrate USP.....1 mg  
Sodium Chloride USP.....3.5 mg  
Sodium Hydroxide/HCl USP.....q.s.  
Water for Injections USP ..... q.s. to 1 ml

1.1 Is this product is licensed to be placed on the market for use in exporting country?

Yes ☒ No ☐ Not applicable ☐

1.2 Is this product naturally on the market in the exporting country? Yes ☒ No ☐ Unknown ☐

(If the answer to 1.2 is yes, continue with Question 2A & omit Question 2B & if answer to 1.2 is No, omit the Question 2A and continue with Question 2B)

2A

1. Product License & date of Issue.  
MB/05/255, 26/02/2021
2. Product License holder (Name and add.)  
United Biotech (P) Limited  
Bagbania, Baddi-Nalagarh Road  
District-Solan (HP) 174101 India
3. Status of applicant a/b/c (key in appropriate Category as define in note)  
a ☒ b ☐ c ☐
4. Permission letter no.  
Is an approved technical summary appended?  
Yes ☐ No ☒ Not provided ☐
5. Is the attached officially approved product Information complete and consonant with the License  
Yes ☐ No ☐ Not provided ☒
6. Applicant for certificate, if different from license holder (name & add.) : SAME

2B

1. Applicant for certificate  
(Name & Address)
2. Status of applicant a/b/c (key in appropriate category as define in note)  
  
a ☐ b ☐ c ☐
3. Why is authorization lacking?  
Not Required ☐  
Not Required ☐  
Under consideration ☐  
Refused ☐
4. Remarks:

3. Does the certifying authority arrange for periodic inspection of manufacturing plant in which the dosage form is produced? <sup>14</sup> Yes ☒ No ☐ Not applicable ☐

3.1 Periodicity of routine inspection: Once in a year.

3.2 Has the manufacturer of this type of dosage forms been inspected? : Yes ☒ No ☐

3.3 Does the facility and operation conform to GMP as recommended by the World Health Organization?

Yes / No / Not applicable      Yes ☒ No ☐ Not applicable ☐

4. Does the information submitted by the applicant satisfy the certifying Authority on all aspects of the manufacturer of the product? Yes ☒ No ☐ if no explain ☐

Address of the certifying authority  
Office of the State Drugs Controller  
Licensing Authority

Health & Family Welfare Department, Himachal Pradesh

Sai Road, Baddi, District Solan, 173205 (H.P.) India

Name of the Authorizing person:

(Dr. Manish Kapoor)

Signature : DEPUTY DRUGS CONTROLLER

-cum-LICENSING AUTHORITY

O/o STATE DRUGS CONTROLLER

BADDI DISTRICT SOLAN, H.P.-173205

E mail ddc4hp@gmail.com

Phone 01795-244288

THIS CERTIFICATE CONFIRMS TO THE FORMAT RECOMMENDED BY THE WORLD HEALTH ORGANIZATION