

GOVERNMENT OF HIMACHAL PRADESH
Health & Family Welfare- Department, Himachal Pradesh
CERTIFICATE OF PHARMACEUTICAL PRODUCTS

No. of Certificate : HFW-H (DRUGS) 427/05/24-134
Valid up to : 21.02.2026

Exporting (certifying) Country: INDIA
Importing (requesting) Country: MONGOLIA

1.0 Proprietary Name (If applicable) and Dosages form of Product : UNIRAUT 100
Active ingredient(s) and amount per unit dose: Iron Sucrose Injection USP 100mg/5ml

Each ml contains:
Ferric Hydroxide complex with Sucrose
eq. to Elemental Iron.....20mg
Water for Injection USP.....q.s.

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, 10/03/2021
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? ¹⁴ 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 certifying authority:

Assistant Drugs Controller
Cum-Licensing Authority
O/o State Drugs Controller
Baddi, Distt. Solan, H.P.173205
sdc4hp@gmail.com, 01795-244288

Name of the Authorizing person: Dr. Kamlesh Naik

Signature :

Stamp & Date (Dr. Kamlesh Naik)



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Cum-Licensing Authority
O/o State Drugs Controller
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18 JUL 2024

THIS CERTIFICATE CONFIRMS TO THE FORMAT RECOMMENDED BY THE WORLD HEALTH ORGANIZATION
(GENERAL INSTRUCTION AND EXPLANATORY NOTES ATTACHED)