Health & Family Welfare Department Himachal Pradesh Baddi, Distt. Solan

Certificate of Good Manufacturing Practices

This one page certificate conforms to the format recommended by the World Health Organization [General Instructions and Explanatory Notes attached].

Certificate No. HFW-H [Drugs] 427/05

On the basis of the inspection carried out on 9th February 2021 and 10th February 2021, we certify that the site indicated on this certificate complies with Good Manufacturing Practices for the dosage forms, categories and activities listed in Table I:

1. Names and Address of Site:

M/s United Biotech (P) Ltd., Bagbania, Baddi-Nalagarh, Road,

Distt. Solan [H.P.]-174101.

2. Manufacturer's License No:

Valid upto 21.02.2026 MNB/05/254 & MB/05/255

3. Table-I:

Dosage Form[s]	Category[ies]	Activity[ies] Production, Packing & Quality Control	
Tablets	General, Betalactam & Oncology		
Capsules (Hard & Soft Gelatin)	General, Betalactam & Oncology	Production, Packing & Quality Control	
Oral Sachet (Powder & Granules)	General	Production, Packing & Quality Control	
Injectables (Liquid, dry & Lyophilized)	General & Oncology	Production, Packing & Quality Control	
Dry Syrups	Betalactam	Production, Packing & Quality Control	
Liquid Orals	General	Production, Packing & Quality Control	
Ointments	General	Production, Packing & Quality Control	
Eye/Ear/Nasal Preparations	General	Production, Packing & Quality Control	
Dry Powder Injections	Betalactam	Production, Packing & Quality Control	
Dry Powder Injections with Diluents	Cephalosporin	Production, Packing & Quality Control	
Soft Gelatin Capsules	General	Production, Packing & Quality Control	

The responsibility for the quality of the individual batches of the pharmaceutical products manufactured through this process lies with the manufacturer.

This certificate remains valid until 21.02.2026. It becomes invalid if the activities and/or categories certified herewith are changed or if the site is no longer considered to be in compliance with GMP.

Address of Certifying Authority: Deputy Drugs Controller

-cum- Licensing Authority O/o State Drugs controller, Baddi, Distt. Solan, H.P.173205 01795-244288,ddc4hp@gmail.com

Name & Function of

(Manish Kapoor)

Responsible person:

Deputy Drugs Controller -cum- Licensing Authority O/o State Durgs Controller, H.P.

01795-244288, ddc4hp@gmail.com

Telephone/Fax No:

Signature:

Date:-

(Dr. Manish Kapoor) 23 27 27 YEPUTY DRILLOS -cum-LICENSING AUTHORITY Stamp:

O/o STATE DRUGS CONTROLLER BADDI DISTRICT SOLAN, H.P-173205

E mail ddc4hp@gmail.com Phone 01795-244088

Explanatory Notes:

- 1. This certificate, which is in the format recommended by WHO certifies the status of the site, listed in point I of the certificate.
- 2. The certificate number should be traceable within the regulatory authority issuing the certificate.
- 3. Where the Regulatory Authority issues a license for the Site, this number should be specified. Record 'Not Applicable" in cases where there is no legal framework for the issuing of a license.
- 4. Table I

List the Dosage Forms, starting materials, categories and activities. Examples are given below:

Example 1

Example 1		
Pharmaceutical	Category [ies]	Activity [ies]
Product[s]1		
Dosage Form [s]:		
Tablets	Cytotoxic	Packaging
	Hormone	Production, Packing, Quality Control
	Penicillin	Repackaging and Labeling
Injectables	Cephalosporin	Aseptic preparation, Packaging, Labeling

Example 2

Pharmaceutical	Category [ies]	Activity [ies]	
Product[s]1	iga rouskorfi i	LECTERIST S.	uid Chala
Starting Material [s]			angeren
Paracetamol	Analgesic	Synthesis, Purification, packing, Labeling	

Use, whenever available, International Non proprietary Names [Inns] or otherwise national Non proprietary Names

- 5. The certificate remains valid until the specified date. The certificate becomes invalid if the activities and/or categories certified are changed or if the site is no longer considered to be in compliance with GMP.
- 6. The requirements for good practices, the manufacture and quality control of drugs referred to in the certificate are those included in Quality Assurance of Pharmaceuticals: a compendium of guidelines and related materials. Good Manufacturing Practices and Inspection. Volume 2, 1999 World Health Organization. Geneva and subsequent updates.

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