Health & Family Welfare Department Himachal Pradesh

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-N/ADC/Drugs/DMLL/2022/127

On the basis of the inspection carried out on 28th & 29th April 2022, 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 Neoveritas Healthcare Private Limited,

At. Mauza Ogli, Suketi Road,

Kala Amb, District Sirmour (H.P.)173030 (India)

2. Manufacturer's License No.:

NL-MB/2022/339 on form 28A

3. Table-I:

Dosage Form[s]	Category[ies]	Activity[ies]
Small Volume Parenterals (Dry)	Cephalosporin and General	Production, Packing & Quality Control
Small Volume Parenterals (Liquid)(Vials & Ampoules)	General	Production, Packing & Quality Control
Ophthalmic Preparations	General	Production, Packing & Quality Control
Small Volume Parenterals (Dry)	Beta Lactum	Production, Packing & Quality Control

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

This certificate now remains valid until **29.05.2025.** 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:

State Drugs Controller,

Controlling cum Licensing Authority

2nd floor, Himuda Commercial Complex, Phase-I,

Housing Board, Baddi, Distt. Solan [H.P.]173205, INDIA.

Name & Function of Responsible person:

Navneet Marwaha

State Drugs Controller,

Controlling cum Licensing Authority, 01795-244288, sdc4hp@gmail.com

Telephone/Fax No: Date: /y/10/2022

Signature: Stamp:

Controlling cum Licensing Author to Baddi Distt. Solan (H. P.)-173205 ju.10.2

11795-244288.sdc4hp@gmail.com

Explanatory Notes:

- 1. This certificate, which is in the format recommended by WHO certifies the status of the site, listed in the 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

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

Example 2

Pharmaceutical Product[s]1	Category [ies]	Activity [ies]
Starting Material[s]		
Paracetamol	Analgesic	Synthesis, Purification, Packing, Labeling

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

- 5. The certificate remains valid until the specified date. The certificate become 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 manufacturer and quality control of drugs referred to in the certificate are those included in Quality Assurance of Pharmaceuticals: a compendium of guidelines are related materials. Good Manufacturing Practices and Inspection. Volume 2, 1999 World Health Organization Geneva and subsequent updates.