

VERNMENT OF THE PEOPLE'S REPUBLIC OF BANGLADESH

MINISTRY OF HEALTH & FAMILY WELFARE DIRECTORATE GENERAL OF DRUG ADMINISTRATION OUSHAD BHABAN, MOHAKHALI DHAKA-1212, BANGLADESH www.dgda.gov.bd



CERTIFICATE OF GOOD MANUFACTURING PRACTICE (GMP) FOR PHARMACEUTICAL (PRODUCTS)

This certificate conforms to the format recommended by the World Health Organization (WHO)

Certificate Number: DA/6-91/04/ 14599

Date: 30-08 It is hereby certified that M/s. Incepta Pharmaceuticals Ltd, a drug (Pharmaceutical Products) manufacturing and marketing organization, has been given license to manufacture and sell its products freely in the People's Republic of Bangladesh as lawfully required and granted in pursuance of The Drugs Act, 1940 (XXIII of 1940) and The Drugs (Control) Ordinance, 1982.

On the basis of inspection carried out on 07-04-2021 & 28-07-2021 we certify that the site indicated on this certificate complies with Good Manufacturing Practices for the dosage forms, categories and activities listed in Table 1.

1. Name & Address of Site:

Incepta Pharmaceuticals Ltd

Dewan Idris Road, Bara Rangamatia, Zirabo, Ashulia, Savar,

Dhaka-1341, Bangladesh

Manufacturer's License No.:

Non-Biological-193

Biological-108

Date of Issue: 27-06-1967 Date of Issue: 27-06-1989

Table: 1

Dosage Form(s) Category(ies) Activity(ies) Antibiotics (Cephalosporins; beta-Tablet (Uncoated, Coated, Procurement of starting and Delayed-Release, Extendedlactam (except Cephalosporins); packaging materials from Release, Controlled Release, Non beta-lactam), Antihypertensive approved sources Modified Release. and Cardioprotectives, Analgesics Quarantine of starting and Sublingual, Buccal, and Antipyretics, Steroids, packaging materials in Cheweble, Effervescent, Antidiarrhoeal, Antacids and warehouse Antiulcerants, Hematinics, Vitamins Orodispersible, MUPS, Sampling and testing of starting and Minerals, Antifungals, Vaginal Tablets), Hard and packaging materials Capsule (Powder Filled, Antidiabetics, Anthelmintics, Storage of released starting and Antiamebics, Antiemetics, Pellet Filled, Liquid Filled), packaging materials Laxatives, Antispasmodics, Powders, Coated Granules, Dispensing of starting and Antihistamines and Antiallergics, Oral Drops, Powder for Oral packaging materials Drops, Nasal drops, Eye Expectorants, Antiasthmatics and Control of manufacturing Drops (Solution, Suspension, Bronchodilators, Anxiolytics, environment Antidepressants, Antimigraine, Emulsion), Nasal Sprays Manufacturing of Finished (Solution, Suspension), Nutrient supplements, Products and Biotherapeutic Gynaecological drugs, Syrup, Injectables (Injections Drug Substances Antipsychotic, Anti-inflamatory, - Intravenous, Intramuscular, In process control of Opthalmics, Antiepileptics, Subcuteneous, Intra-Intermediates, Bulks and Anticonvulsants, Anticoagulants, Articular, Lyophilized Finished Products and Antiplatelets, Diuretics, Insulins, Injections, Powder for Biotherapeutic Drug Substances Injections, Intravenous Lipid lowering agents, Medication Infusions), for Hepatitis B and C, Antivirals,

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Dosage Form(s)	Category(ies)	Activity(ies)
Oral solution, Powder for Oral solution, Nebulized Solution, Nebulizer Suspension, Oral suspension, Oral Emulsion, Powder for Oral Suspension, Elixir; Respiratory Solution, Dry Powder for Inhalation, Biotherapeutic Drug Product, Biotherapeutic Drug Substance	Immunosuppressants, Anesthetics, Antifibrinolytics, Antimalarial, Tinibs, Monoclonal Antibody (mAbs), Cytokines, Hematopoietic Agents, Antigout, Urologicals, Tyrosine Kinase Inhibitors, CDK 4/6 Inhibitor, Antirheumatic.	 Control of packaging and labeling Quality Assurance of finished products and biotherapeutic drug substances Storage of finished products Effluent treatment and safe disposal of pharmaceutical wastes Stability studies

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

The manufacturing plant in which the pharmaceutical products are produced is subject to inspection at suitable intervals.

The manufacturer conforms to the requirements for good practices in the manufacture and quality control (GMP) of drugs, as required under law in this country, as well as recommended by the World Health Organization (WHO) in respect of pharmaceutical products to be manufactured, sold or distributed within the country of origin or to be exported.

This certificate remains valid for a period of 2 (Two) years from the date of issue. 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.

Name of Authorized Person

: Major General Mohammad Yousuf Address of the Certifying Authority : Directorate General of Drug Administration

Mohakhali, Dhaka-1212

Telephone Fax No E-mail Web-site

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Stamp and Date:



Major General Mohammad Yousuf Director General

Directorate General of Drug Administration

Licensing Authority (Drugs)

Govt. of the People's Republic of Bangladesh