महाराष्ट्र शासन आयुक्त अन्न वा औषध प्रशासन , महा. राज्य ३४१ , वांद्रे - कुर्ला संकुल , रिजर्व बँक समोर, वांद्रे (पूर्व) मुंबई - ४०० ०५१.



GOVERNMENT OF MAHARASHTRA

COMMISSIONER

Food and Drugs Administration (M.S.) 341, Bandra-Kurla Complex, Opposite of RBI Buildings, BAndra (E), Mumbai - 400 051 Tel: 022 - 26592362-65

E-Mail: comm.fda-mah@nic.in

ず. NEW-WHO-GMP/CERT/KD/74763/2018/ <u>2090</u> /11

दिनांक. 04/07/2018

प्रति, PROTO CHEMICAL INDUSTRIES PALGHAR

विषय - डब्लूएचओ - जीएमपी प्रमाणपत्र मंजुरीबाबत

संदर्भ - आपला प्रस्ताव क्रमांक 74763

महोदय,

सोबत डब्लूएचओ - जीएमपी प्रमाणपत्र / सीओपीपी (सर्टिफिकेट ऑफ फार्मास्युटिकल्स प्रॉडक्ट्स / स्टेटमेंट ऑफ लायसन्सिंग) स्टेटस प्रमाणपत्र क्रमांक डब्लूएचओ - जीएमपी/ KD/74763 (एकूण प्रमाणपत्रे 1) पाठवीण्यात येत आहेत.

सहआयुक्त, मुख्यालय, मुंबई यांचे मान्यतेने

(सु सं मोहिते)

सहायक आयुक्त (मुख्यालय) का क्र.११ अन्न व औषध प्रशासन , म. राज्य.



Office of The Commissioner, Food & Drugs Administration M.S. Bandra – Kurla Complex, Bandra (E), Mumbai – 400 051

Date: 04/07/2018

CERTIFICATE OF GOOD MANUFACTURING PRACTICES

This Certificate conforms to the format recommended by the World Health Organization. (General instructions and explanatory notes attached).

Certificate No.: NEW-WHO-GMP/CERT/KD/74763/2018/11/24067

On the basis of the inspection carried out on **05/06/2018**, **06/06/2018** and **29/06/2018**, 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.

Name of the Firm

PROTO CHEMICAL INDUSTRIES

Address

PLOT NO. E-121,MIDC TARAPUR BOISAR PALGHAR 401506 MAHARASHTRA STATE,

INDIA

2. Licence No.

KD778 In Form 25

Table 1

Sr.No.	Dosage Form(s)	Categor(ies)	Activity(ies)	
	Active Pharmaceutical Ingredients (Bulk Drugs)	General (Other than Cephalosporins, Penicillin, Cytotoxic, Hormones)	Production, Filling, Packing, labelling, Quality Control, Quality Assurance	

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 02 Jul 2021 . 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: Food & Drug Administration, M.S.

Bandra-kurla Complex, Bandra (E), Mumbai – 400 051.

Maharashtra, INDIA. Tel: +91-22-26592363/64

Fax: +91-22-26591959 10RP0857476320180703 Name of the Authorised person : A. T. NIKHADE

Signature:

Stamp and Date : Joint Commissioner (HQ) & Controlling

Authority

Food & Drug Administration, M.S.

Bandra (E), Mumbai. Maharashtra State, India

Date: 03 Jul 2018

0 3 JUL 2018

Explanatory notes

- 1. This certificate which is in the format recommended by WHO, certifies the status of the site listed in point 1 of the certificate.
- 2. The certification number should be traceable within the regulatory authority issuing the certificate.
- 3. Where the regulatory authority issues a licence for the site, this number should be specified record "not applicable" in cases where there is no legal framework for the issuing of a licence.
- 4. Table 1
 List the dosage forms, starting materials, categories and activities. Examples are given below.

Example -1

Pharmaceutical Product (s)1	Category (ies)	Activity (ies)	
Dosage form (s)			
Tablets	Cytotoxic	Packaging	
	Hormone	Production, Packaging, Quality control.	
Injectables	Penicillin	Repackaging & Labelling.	
	Cefalosporin	Aseptic preparation, Packaging, Labelling.	

Example - 2.

Pharmaceutical Product (s)1	Category (ies)	Activity (ies)
Starting material (s)2		
Paracetamol	Analgesic	Synthesis, Purification, Packing, Labelling.

Use, whenever available. International Nonproprietary Names (INNs) or otherwise national nonproprietary names.

- 5. The certificate remains valid until the specified date. The certificate becomes invalid if the activities and/or categories certified are changed or it the site and 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.

HIDS III ON

LIST OF PRODUCT APPROVED UNDER WHO GMP¹

No. of certificate

NEW-WHO-GMP/CERT/KD/74763/2018/11

VALID UP TO :02 Jul 2021

/24067

Name of Manufactring Firm

PROTO CHEMICAL INDUSTRIES

PLOT NO. E-121, MIDC TARAPUR BOISAR

PALGHAR 401506 MAHARASHTRA STATE, INDIA

Drug License No

KD778 In Form 25

Sr.No.	Name of the Product		Composition	
1	POVIDONE IODINE BP	Povidone Iodine BP		
2	POVIDONE IODINE EP	Povidone Iodine EP		b
3	POVIDONE IODINE IP	Povidone Iodine IP		
4	POVIDONE IODINE USP	Povidone Iodine USP		

Address of certifying authority: Food & Drug Administration, M.S. Bandra-kurla Complex,

Bandra (E), Mumbai – 400 051.

Maharashtra,INDIA. Tel: +91-22-26592363/64 Fax: +91-22-26591959

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Name of the Authorised person : A. T. NIKHADE

Signature:

Stamp and Date : Joint Commissioner (HQ) & Controlling Authority

Food & Drug Administration, M.S.

Bandra (E), Mumbai. Maharashtra State, India Date:03 Jul 2018

0 3 JUL 2018