



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

13 AUG 2018

INSTITUTO DE SALUD PUBLICA
Certifico que la presente
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que he tenido a la vista
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páginas
Stgo. 11-01-19

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/72365/2018/11/24339

On the basis of the inspection carried out on 07/06/18, 08/07/18 and 18/07/2018, 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.

1. Name of the Firm : CIRON DRUGS & PHARMACEUTICALS PVT. LTD.
Address : N-118, 118/1, 119, 119/1, 119/2, 113 MIDC, TARAPUR, BOISAR, DIST. THANE 401506 MAHARASHTRA STATE, INDIA
2. Licence No. : KD80 In Form 25, KD74 In Form 28, KD/3 In Form 28B

Table 1

Sr.No.	Dosage Form(s)	Categor(ies)	Activity(ies)
1	External Preparation (Ointments / Creams / Lotion/Gel/Ear drop/Nasal drop/Spray)	General (Other than Cephalosporins, Penicillin, Cytotoxic, Hormones)	Production, Filling, Packing, labelling, Quality Control, Quality Assurance
2	Eye / Ear Drops	General (Other than Cephalosporins, Penicillin, Cytotoxic, Hormones)	Production, Filling, Packing, labelling, Quality Control, Quality Assurance
3	Eye Drops / Ophthalmic Preparations	General (Other than Cephalosporins, Penicillin, Cytotoxic, Hormones)	Production, Filling, Packing, labelling, Quality Control, Quality Assurance
4	Inhalation	General (Other than Cephalosporins, Penicillin, Cytotoxic, Hormones)	Production, Filling, Packing, labelling, Quality Control, Quality Assurance
5	Liquid Injection (SVP)	General (Other than Cephalosporins, Penicillin, Cytotoxic, Hormones)	Production, Filling, Packing, labelling, Quality Control, Quality Assurance
6	Liquid Orals	General (Other than Cephalosporins, Penicillin, Cytotoxic, Hormones)	Production, Filling, Packing, labelling, Quality Control, Quality Assurance
12			

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 01 Aug 2021. It becomes invalid if the activities and / or categories certified hereon 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: 01-22365933334
Fax: 01-2236591956

Name of the Authorised person : A. T. NIKHADE

Signature :

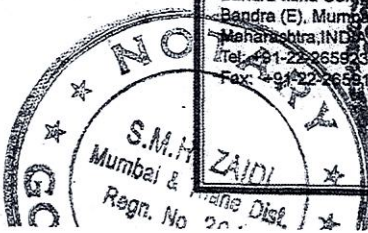
Stamp and Date : Joint Commissioner (HQ) & Controlling Authority
Food & Drug Administration, M.S.
Bandra (E), Mumbai.
Maharashtra State, India
Date: 09 Aug 2018

A copy of this document / CERTIFICATE has been recorded with the Chamber

MR. SAMIR PAUL PINTO
ASSISTANT MANAGER

Authorized Signatory

Bombay Chamber of Commerce and Industry
Regn. No. 105872



09 AUG 2018



भारत सरकार / GOVERNMENT OF INDIA
अपोस्टिल / APOSTILLE
(Convention de La Haye du 5 octobre 1961)
INDIA

This public document of the type
COMMERCIAL DOCUMENT

is issued to CIRON DRUGS & PHARMACEUTICALS
PVT. LTD.

has been signed by SAMIR PAUL PINTO

with the seal / stamp of ASSTT. MANAGER, BOMBAY CHAMBER
OF COMMERCE AND INDUSTRY

Certified by
Section Officer(OI) MINISTRY OF EXTERNAL AFFAIRS
on 27-Nov-2018 at NEW DELHI, INDIA

with reference no. MHMC0035552818

Seal / Stamp

Signature

(SABARATHAN PAUL)
Section Officer (O.I.)
विदेशी मामलों, नई दिल्ली
Ministry of External Affairs
New Delhi





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2. Licence No. : **KD80 In Form 25, KD74 In Form 28, KD/3 In Form 28B**

Table 1

Sr.No.	Dosage Form(s)	Categor(ies)	Activity(ies)
7	Lyophilised / Powder injectable	General (Other than Cephalosporins, Penicillin, Cytotoxic, Hormones)	Production, Filling, Packing, labelling, Quality Control, Quality Assurance
12			

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Tel: +91-22-26592363/64
Fax: +91-22-26591959

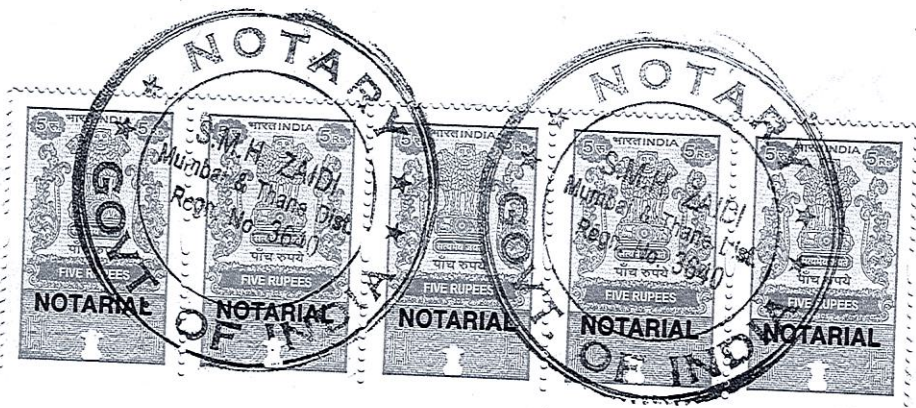
Name of the Authorised person : **A. T. NIKHADE**

Signature :

Stamp and Date : **Joint Commissioner (HQ) & Controlling Authority**
Food & Drug Administration, M.S.
Bandra (E), Mumbai.
Maharashtra State, India
Date: 09 Aug 2018



15 NOV 2018



ATTESTED BY ME


S. M. H. ZAIDI
NOTARY

Government of India
Mumbai & Thane Dist

15 NOV 2018