

**FOOD SAFETY & DRUGS ADMINISTRATION, AUTHORITY
DIRECTORATE GENERAL OF MEDICAL HEALTH AND FAMILY
WELFARE, SAHASTRADHARA ROAD, DEHRADUN**

F.No. 17P/1/71/2006/ 27142

Date: 10-12-2019

Certificate of Good Manufacturing Practices

Certificate No.: 17P/1/71/2006 / 27142

- 1- On the basis of the Joint Inspection carried out on 20-11-2019 & 22-11-2019, and Further inspected by I.D HQ on Dated 09-12-2019 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.
- 2- **Name & Address of site:**
M/s Jubilant Generics Ltd. (Formerly Jubilant Lifesciences Ltd.)
Village Sikandarpur Bhainswal,
Roorkee- Dehradun Highway, Bhagwanpur,
Roorkee, Distt. Haridwar. Uttarakhand.
- 3- **Manufacturer's license number:**
Form 25- 58/UA/2007
Form 28- 81/UA/SC/P-2010

4- Table 1:

Dosage form(s)	Activity(ies)
Tablets (Non Beta Lactum)	Manufacturing
Capsules (Non Beta Lactum)	Manufacturing

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 09-12-2022. 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.

The firm is following Good Manufacturing Practices as per World Health Organization (WHO) TRS Guide Lines, in the Manufacturing & testing of the said categories of Products and Items in respect of which the Certificates of Pharmaceuticals products have been issued.

Address of certifying Authority:

Directorate General of Medical Health & Family Welfare,
Sahastradhara Road, Dehradun (Uttarakhand) INDIA.

Name & function of responsible person:

Shri Tajber Singh
Drugs Licensing & Controlling Authority
Uttarakhand.

Email: drugcontroluk@gmail.com

Tel. no. NA

Fax. no. 0135260874



**NOTARY PUBLIC
DELHI (INDIA)**

4 MAR 2020



(Tajber Singh)

**Drugs Controlling & Licensing Authority
(Uttarakhand)
(Tajber Singh)
Drug Controlling & Licensing Authority (Mfg.)
(Uttarakhand)**



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

Country

REPUBLIC OF INDIA

This public document

COMMERCIAL DOCUMENT

has been signed by DIRECTOR

acting in the capacity of DIRECTOR

bears the seal/stamp of PHD CHAMBER OF COMMERCE AND
INDUSTRY, NEW DELHI

Certified

at NEW DELHI, INDIA the 11-Mar-2020

by SO (OI/Attestation) MINISTRY OF EXTERNAL AFFAIRS

No. DLND0007249420

Seal / Stamp

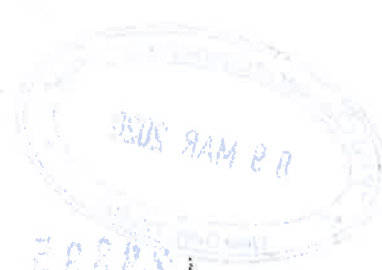
is issued to JUBILANT GENERICS LTD.

Signature



(सुनील चनाप)
(SUNIL CHANAP)
अध्यापक (ओ.आई.)
Section Officer (OI)
नो. वी. वी. प्रभाग / C.P.V. Div.
विदेश मंत्रालय, नई दिल्ली
Ministry of External Affairs, New

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Attested
New Delhi (India)
Ministry of External Affairs
New Delhi (India)

**SEGURIDAD DE ALIMENTOS & ADMINISTRACIÓN DE MEDICAMENTOS, DIRECCIÓN
GENERAL DE SALUD MÉDICA Y BIENESTAR FAMILIAR,
SAHASTRADHARA ROAD, DEHRADUN**

F. No. 17P/1/71/2006/27142

Fecha: 10-12-2019

Certificado de Buenas Prácticas de Manufactura

Certificado No.: 17P/1/71/2006/27142

1- En base a la Inspección Conjunta llevada a cabo el 20-11-2019 & 22-11-2019 y adicionalmente inspeccionada por I.D HQ con fecha 09-12-2019, certificamos que el sitio indicado en este certificado cumple con las Buenas Prácticas de Manufactura para las formas de dosis, categorías y actividades listadas en la Tabla 1.

2- Nombre & Dirección del sitio:

M/s Jubilant Generics Ltd. (Anteriormente Jubilant Lifesciences Ltd.)
Village Sikandarpur Bhainswal,
Roorkee- Dehradun Highway, Bhagwanpur,
Roorkee, Distt. Haridwar. Uttarakhand.

3- Número de licencia de fabricante:

Forma 25 – 58/UA/2007
Forma 28 – 81/UA/SC/P-2010

Traducción fiel al original

4- Tabla 1:

Forma(s) de Dosis	Actividad(es)
Tabletas (No Betalactámico)	Fabricación
Cápsulas (No Betalactámico)	Fabricación

La responsabilidad por la calidad de los lotes individuales de productos farmacéuticos fabricados a través de este proceso yace con el fabricante.

Este certificado permanece válido hasta el 09-12-2022. Queda inválido si las actividades y/o categorías certificadas en el presente son cambiadas o si el sitio deja de ser considerado como en cumplimiento con las BPM.

La firma está siguiendo las **Buenas Prácticas de Manufactura según las Directrices TRS de la Organización Mundial de la Salud (OMS)**, en la Fabricación & prueba de dichas categorías de Productos y Artículos con respecto de los cuales el Certificado de Productos Farmacéuticos ha sido emitido.

Dirección de la Autoridad certificante:

Dirección General de Salud Médica & Bienestar Familiar,
Sahastradhara Road, Dehradun (Uttarakhand) INDIA.

Nombre & función de la persona responsable:

Shri Tajber Singh
Autoridad Controladora & Licenciadora de Medicamentos
Uttarakhand

Email: drugcontroluk@gmail.com

Tel. no. NA

Fax no. 0135260874



(Tajber Singh)

**Autoridad Controladora & Licenciadora de Medicamentos
(Uttarakhand)**

(Tajber Singh)

Drug Controlling & Licensing Authority (Mfg.)
(Uttarakhand)

NOTARY PUBLIC
DELHI (INDIA)

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 No. DLND0007249420

Seal / Stamp



Issued to JUBILANT GENERICS LTD.

Signature

(सुनील चनाप)
 (SUNIL CHANAP)
 अधिवक्ता (ओआई) (सी.ओ.ओ.)
 Section Officer (OI)
 श्री. ए. वी. चनाप / C.P.V. Divya
 विदेश सहायता, नई दिल्ली
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For Signatory

Ministry of External Affairs, New Delhi