

FOOD AND DRUGS AUTHORITY

GMP CERTIFICATE

The Food and Drugs Authority, Ghana, after an Audit conducted in <u>APRIL</u> . 2017.
certifies that the companySYNOKEM PHARMACEUTICAL LIMITED.
with manufacturing site at PLOT.35-36, SECTOR-6A, I.I.E. (SIDCUL) RANIPUR (BHELL),
HARIDWAR - 249 403

is able to maintain an acceptable standard of Good Manufacturing Practices (**GMP**) as per the guidelines of the World Health Organisation on current codes of GMP and conforms with **Section 131 of the Public Health Act**, **2012, Act 851 of the Republic of Ghana.**

The Authority hereby authorizes the company to manufacture the following pharmaceutical dosage forms:

1. ORAL TABLETS	4. HORMONES (TABS / CAPSULES)
2. ORAL CAPSULES	5. XX
3. ORAL SUSPENSIONS AND SYRUPS	6. XX

which were included in the afore stated audit for supplies to the Republic of Ghana. This certificate must be reproduced in full to interested parties upon request.

Certificate No.: ...FDA/GMP/004/06/17

Expiry Date: MARCH, 2022

DELESE A. A. DARKO (MRS) (AG. CHIEF EXECUTIVE OFFICER)

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This certificate remains the property of Food and Drugs Authority, Ghana and must be returned upon written demand. Any alteration to this certificate renders it null and void



Food and Drugs Authority

Head Office P. O. Box CT 2783, Cantonments, Accra-Ghana Tel: (+233-302)233200, 235100 Fax: (+233-302)229794, 225502 Email: fda@fdaghana.gov.gh

FDA/DRID/DED/PIU/17/0188

30th June 2017

The Managing Director Synokem Pharmaceuticals Ltd Synokem House 14/486 Sunder Vihar Paschim Vihn New Delhi 110097 India

Dear Sir/Madam,

GOOD MANUFACTURING PRACTICE (GMP) CERTIFICATION

Pursuant to the provisions of **Section 131 of the Public Health Act, 2012, Act 851 of the Republic of Ghana,** this is to inform you that the Food and Drugs Authority, Ghana, after conducting Good Manufacturing Practices (GMP) audit at your facility sited at **Haridwar, India**, during the period of 25th- 26th May, 2017, has determined that the operations of the facility meet acceptable GMP standard. The facility is therefore licensed to manufacture dosage forms as covered in the certification for the Ghanaian market.

The Food and Drugs Authority hereby issues you with a GMP certificate with number: **FDA/GMP/004/06/17**, valid up to **April**, 2022 and subject to renewal.

You are directed to act in accordance with the under listed terms and conditions of the certification.

- 1. The certificate should be renewed after its expiry.
- 2. Only product(s) manufactured in the audited facility can be distributed on the Ghanaian market.
- 3. Distribution of the product(s) covered by the certification in the Ghanaian market ceases after the expiration of the certification.

Yours faithfully,

and

DELESE A. A. DARKO (MRS) AG. CHIEF EXECUTIVE OFFICER

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